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(54) Title: NODULISPORIC ACID DERIVATIVES

126 East Lincoln Avenue, Rahway, NJ 07065 (US).

(57) Abstract

The present invention relates to novel nodulisporic acid derivatives, which are acaricidal, antiparasitic, insecticidal and anthelmintic agents.

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TITLE OF THE INVENTION NODULISPORIC ACID DERIVATIVES

CROSS REFERENCE

This is a continuation-in part of co-pending application U.S.S.N. 08/406,619, filed March 20, 1995, which is hereby incorporated by reference.

BACKGROUND OF THE INVENTION

Nodulosporic acid and two related components are antiparasitic agents and ectoparasiticidal agents isolated from the fermentation culture of *Nodulisporium* sp. MF-5954 (ATCC 74245). These three compounds have the following structures:

29,30-dihydro-20,30-oxa-nodulisporic acid (compound B)

31-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-nodulisporic acid (compound C)

5 SUMMARY OF THE INVENTION

This invention relates to new acaricidal, antiparasitic, insecticidal and anthelmintic agents related to the nodulisporic acids, to processes for their preparation, compositions thereof, their use in the treatment of parasitic infections, including helminthiasis, in human and animals, and their use in the treatment of parasitic infections in plants or plant products.

DETAILED DESCRIPTION OF THE INVENTION

The present invention provides compounds having the

15 formula I:

10

20 wherein

R₁ is

- (1) hydrogen,
- (2) optionally substituted C₁-C₁₀ alkyl,
- (3) optionally substituted C2-C10 alkenyl,

		(4)	option	ally substituted C ₂ -C ₁₀ alkynyl,
		(5)	option	ally substituted C3-C8 cycloalkyl,
		(6)	option	ally substituted C5-C8 cycloalkenyl
		wher	e the substi	tutents on the alkyl, alkenyl, alkynyl,
5		cyclo	alkyl and c	ycloalkenyl are 1 to 3 groups independently
		selec	ted from	
			(i) C ₁ -	C5 alkyl,
			(ii)	X-C ₁ -C ₁₀ alkyl, where X is O or S(O) _m
			(iii)	C3-C8 cycloalkyl,
10			(iv)	hydroxy,
			(v)	halogen,
			(vi)	cyano,
			(vii)	carboxy,
			(viii)	- , I will I will
15		indep	endently H	or C ₁ -C ₁₀ alkyl,
			(ix)	C ₁ -C ₁₀ alkanoylamino, and
			(x)	aroyl amino wherein said aroyl is
		option	nally substit	tuted with 1 to 3 groups independently
20			ed from Rf	
20		(7)	aryl Co	-C5 alkyl wherein said aryl is optionally
		substi	tuted with 1	to 3 groups independently selected from
		Rf,		
		(8)		perfluoroalkyl
05		(9)	a 5- or 6	5-membered heterocycle containing from 1
25		to 4 he	eteroatoms	independently selected from oxygen, sulfur
		and ni	trogen aton	ns optionally substituted by 1 to 3 groups
		indepe	endently sel	ected from hydroxy, oxo, C1-C10 alkyl
				which may be saturated or partly
30	Do Do o	unsatu	•	1 000 000 1
30				ntly ORa, OCO2Rb, OC(O)NRcRd; or
				or = $N-NRcRd$;
	R5 and R			
	R7 is		represent -	
	17 / 18	(1)	CHO, or	

		(12)
		(13) optionally substituted C3-C8 cycloalkyl
		(14) optionally substituted C5-C8 cycloalkenyl
		where the substituents on the alkyl, alkenyl, alkynyl,
_		alkanoyl, alkenoyl, alkynoyl, aroyl, aryl, cycloalkanoyl,
5		cycloalkenoyl, alkylsulfonyl, cycloalkyl and cycloalkenyl
		are from 1 to 10 groups independently selected from
		hydroxy, C1-C6 alkoxy, C3-C7 cycloalkyl, aryl C1-C3
		alkoxy, NRgRh, CO2Rb, CONRCRd and halogen,
		(15) C ₁ -C ₅ perfluoroalkyl,
10		(16) arylsulfonyl optionally substituted with 1 to 3
		groups independently selected from C1-C5 alkyl, C1-C5
		perfluoroalkyl, nitro, halogen and cyano,
		(17) a 5- or 6-membered heterocycle containing 1 to 4
		heteroatoms selected from oxygen, sulfur and nitrogen
15		optionally substituted by 1 to 4 groups independently
		selected from C ₁ -C ₅ alkyl, C ₁ -C ₅ alkenyl, C ₁ -C ₅
		perfluoroalkyl, amino, C(O)NRcRd, cyano, CO2Rb and
	•	halogen, and which may be saturated or partly unsaturated;
	Rb is	(1) H,
20		(2) optionally substituted aryl,
		(3) optionally substituted C ₁ -C ₁₀ alkyl,
		(4) optionally substituted C3-C10 alkenyl,
		(5) optionally substituted C3-C10 alkynyl,
		(6) optionally substituted C3-C15 cycloalkyl,
25		(7) optionally substituted C5-C10 cycloalkenyl, or
		(8) optionally substituted 5- to 10-membered
		heterocycle containing from 1 to 4 heteroatoms
		independently selected from oxygen, sulfur and nitrogen:
		where the substituents on the aryl, alkyl, alkenyl, cycloalkyl,
30		cycloalkenyl, heterocycle, or alkynyl are from 1 to 10 groups
		independently selected from
		(i) hydroxy,
		(ii) C ₁ -C ₆ alkyl,
		Z1115

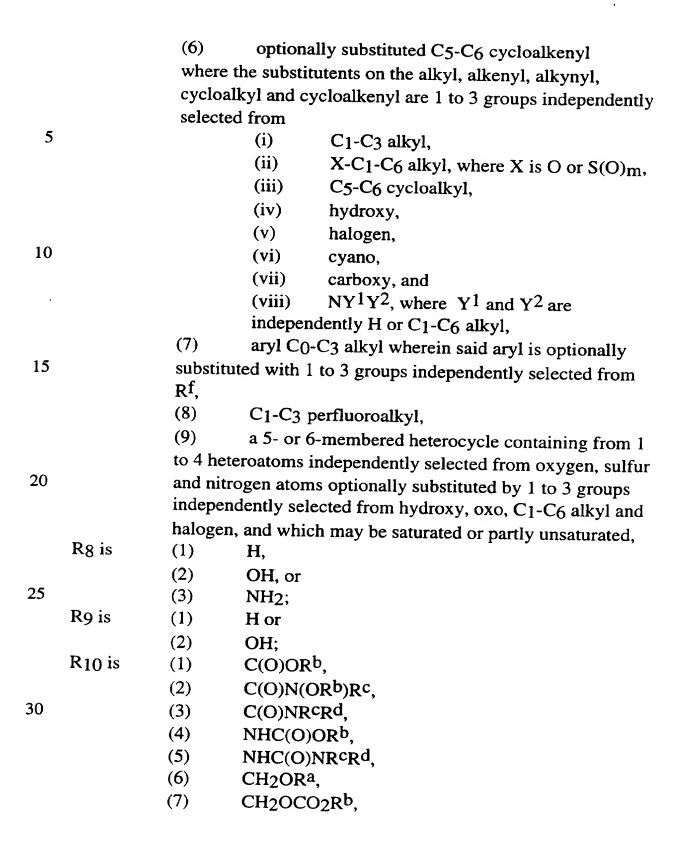
(iii)

oxo,

	(iv)	SO ₂ NRgRh,
	(v)	aryl C1-C6 alkoxy,
	(vi)	hydroxy C ₁ -C ₆ alkyl,
	(vii)	C1-C12 alkoxy,
5	(viii)	hydroxy C ₁ -C ₆ alkoxy,
	(ix)	amino C ₁ -C ₆ alkoxy,
	(x)	cyano,
	(xi)	mercapto,
	(xii)	C ₁ -C ₆ alkyl-S(O)m,
10	(xiii)	C3-C7 cycloalkyl optionally substituted
	with 1 to	o 4 groups independently selected from Re
	(xiv)	C5-C7 cycloalkenyl,
	(xv)	halogen,
	(xvi)	C ₁ -C ₅ alkanoyloxy,
15	(xvii)	C(O)NRgRh,
	(xviii)	CO ₂ R ⁱ ,
	(xix)	formyl,
	(xx)	-NRgRh,
	(xxi)	winen may
20	be saturated or par	rtially unsaturated, containing from 1 to 4
	heteroatoms indep	pendently selected from oxygen, sulfur and
	nitrogen, and optic	onally substituted with 1 to 5 groups
	independently sele	ected from Re,
	(xxii)	optionally substituted aryl, wherein the
25	aryl substituents a	re 1,2-methylenedioxy or 1 to 5 groups
	independently sele	
	(xxiii)	i juliant my ci cy untoxy,
	wherein the aryl si	ubstituents are 1,2-methylenedioxy or 1 to
		lently selected from Re, and
30	(xxiv)	C1-C5 perfluoroalkyl;
	R ^c and R ^d are independently se	elected from Rb; or
	R ^c and R ^d together with the N	to which they are attached form a 3- to 10-
	membered ring con	ntaining 0 to 2 additional heteroatoms
	selected from O, S	(O) _m , and N, optionally substituted with 1

		to 3 gro	oups independently selected from Rg, hydroxy, thioxo
		and oxo	o;
	Re is	(1)	halogen,
		(2)	C ₁ -C ₇ alkyl,
5		(3)	C1-C3 perfluoroalkyl,
		(4)	$-S(O)_{\mathbf{m}}R^{\mathbf{i}}$
		(5)	cyano,
		(6)	nitro,
		(7)	$R^{i}O(CH_{2})_{V^{-}}$
10		(8)	$R^{i}CO_{2}(CH_{2})_{V^{-}},$
		(9)	R ⁱ OCO(CH ₂) _v ,
		(10)	optionally substituted aryl where the substituents
		are from	1 to 3 of halogen, C1-C6 alkyl, C1-C6 alkoxy, or
		hydroxy	',
15		(11)	SO ₂ NRgR ^h , or
	c	(12)	amino;
	Rf is	(1)	C ₁ -C ₄ alkyl,
		(2)	X-C ₁ -C ₄ alkyl, where X is O or $S(O)_m$,
		(3)	C2-C4 alkenyl,
20		(4)	C2-C4 alkynyl,
		(5)	C ₁ -C ₃ -perfluoroalkyl,
		(6)	NY^1Y^2 , where Y^1 and Y^2 are independently H or
		C ₁ -C ₅ a	lkyl,
		(7)	hydroxy,
25		(8)	halogen, and
	_	(9)	C ₁ -C ₅ alkanoyl amino,
	Rg and Rh	are indepe	ndently
		(1)	hydrogen,
		(2)	C ₁ -C ₆ alkyl optionally substituted with hydroxy,
30		amino, o	r CO ₂ R ⁱ
		(3)	aryl optionally substituted with halogen, 1,2-
		methylen	edioxy, C1-C7 alkoxy, C1-C7 alkyl or C1-C3
		perfluoro	

	•	(4)	aryl C1-C6 alkyl, wherein the aryl is optionally
		substi	tuted with C1-C3 perfluorolkyl or 1,2-methylenedioxy;
		(5)	C ₁ -C ₅ alkoxycarbonyl,
		(6)	C ₁ -C ₅ alkanoyl,
5		(7)	C1-C5 alkanoyl C1-C6 alkyl,
		(9)	aryl C1-C5 alkoxycarbonyl,
		(10)	aminocarbonyl,
		(11)	C1-C5 monoalkylaminocarbonyl
		(12)	C ₁ -C ₅ dialkylaminocarbonyl; or
10	Rg and Rl	ogether	with the N to which they are attached form a 3- to 7-
		memb	ered ring containing 0 to 2 additional heteroatoms
		selecte	ed from O, S(O) _m , and N, optionally substituted with 1
	_	to 3 gr	oups independently selected from Re and oxo;
	R ⁱ is	(1)	hydrogen,
15		(2)	C1-C3 perfluoroalkyl,
		(3)	C1-C6 alkyl,
		(4)	optionally substituted aryl Co-C6 alkyl, where the
		aryl su	bstituents are from 1 to 3 groups independently
		selecte	d from halogen, C1-C6 alkyl, C1-C6 alkoxy, and
20		hydrox	y;
	m is	0 to 2;	and
	v is	0 to 3;	
	a pharmace	eutically a	acceptable salt thereof; and
	excluding i	nodulispo	oric acid, 29,30-dihydro-20,30-oxa-nodulisporic acid,
25		and 31-	-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-
		nodulis	poric acid.
		In a pre	sfarred amhadiment the survey of
	compounds	of Form	eferred embodiment, the present invention provides ula I wherein
30	R ₁ is	(1)	hydrogen,
		(2)	optionally substituted C ₁ -C ₆ alkyl,
		(3)	optionally substituted C2-C6 alkenyl,
		(4)	optionally substituted C ₂ -C ₆ alkenyl,
		(5)	optionally substituted C ₂ -C ₆ alkynyl, optionally substituted C ₅ -C ₆ cycloalkyl,
		\- /	-r



		(8)	CH2OC(O)NRCRa,
		(9)	C(O)NRcNRcRd, or
		(10)	C(O)NRcSO2Rb;
	Ra is	(1)	hydrogen,
5		(2)	optionally substituted C1-C6 alkyl,
		(3)	optionally substituted C3-C6 alkenyl,
		(4)	optionally substituted C3-C6 alkynyl,
		(5)	optionally substituted C1-C6 alkanoyl,
		(6)	optionally substituted C3-C6 alkenoyl,
10		(7)	optionally substituted C3-C6 alkynoyl,
		(8)	optionally substituted aroyl,
		(9)	optionally substituted aryl,
		(10)	optionally substituted C5-C6 cycloalkanoyl,
		(11)	optionally substituted C5-C6 cycloalkenoyl,
15		(12)	optionally substituted C1-C6 alkylsulfonyl
		(13)	optionally substituted C5-C6 cycloalkyl
		(14)	optionally substituted C5-C6 cycloalkenyl
		where	the substituents on the alkyl, alkenyl, alkynyl,
		alkano	yl, alkenoyl, alkynoyl, aroyl, aryl, cycloalkanoyl,
20		cycloal	kenoyl, alkylsulfonyl, cycloalkyl and cycloalkenyl
		are from	m 1 to 10 groups independently selected from
		hydrox	y, C1-C4 alkoxy, C5-C6 cycloalkyl, aryl C1-C3
		alkoxy,	NRgRh, CO2Rb, CONRCRd and halogen,
		(15)	C ₁ -C ₃ perfluoroalkyl,
25		(16)	arylsulfonyl optionally substituted with 1 to 3
			independently selected from C1-C3 alkyl, C1-C3
		perfluo	roalkyl, halogen and cyano,
		(17)	a 5- or 6-membered heterocycle containing 1 to 4
		heteroa	toms selected from oxygen, sulfur and nitrogen
30		optiona	lly substituted by 1 to 4 groups independently
		selected	from C1-C3 alkyl, C1-C3 alkenyl, C1-C3
			roalkyl, amino, C(O)NR ^c R ^d , cyano, CO ₂ R ^b and
	_ L .		, and which may be saturated or partly unsaturated;
	Rb is	(1)	Н,

	(2) optionally substituted aryl,
	(3) optionally substituted C ₁ -C ₇ alkyl,
	(4) optionally substituted C3-C7 alkenyl,
	(5) optionally substituted C3-C7 alkynyl,
5	(6) optionally substituted C5-C7 cycloalkyl,
	(7) optionally substituted C5-C7 cycloalkenyl, or
	(8) optionally substituted 5- to 10-membered
	heterocycle containing from 1 to 4 heteroatoms
	independently selected from oxygen, sulfur and nitrogen;
10	where the substituents on the aryl, alkyl, alkenyl, cycloalkyl,
	cycloalkenyl, heterocycle, or alkynyl are from 1 to 10 groups
	independently selected from
	(i) hydroxy,
	(ii) C ₁ -C ₃ alkyl,
15	(iii) oxo,
	(iv) SO ₂ NRgRh,
	(v) aryl C ₁ -C ₃ alkoxy,
	(vi) hydroxy C ₁ -C ₃ alkyl,
	(vii) C ₁ -C ₇ alkoxy,
20	(viii) hydroxy C ₁ -C ₃ alkoxy,
	(ix) amino C ₁ -C ₃ alkoxy,
	(x) cyano,
	(xi) C ₁ -C ₃ perfluoroalkyl,
	(xii) C ₁ -C ₃ alkyl-S(O)m,
25	(xiii) C5-C6 cycloalkyl optionally substituted
	with 1 to 4 groups independently selected from Re,
	(xiv) C5-C6 cycloalkenyl,
	(xv) halogen,
	(xvi) C ₁ -C ₃ alkanoyloxy,
30	(xvii) C(O)NRgRh,
	(xviii) CO ₂ R ⁱ ,
	(xix) optionally substituted aryl C1-C3 alkoxy,
	wherein the aryl substituents are 1,2-methylenedioxy or 1 to
	5 groups independently selected from De

5 groups independently selected from Re,

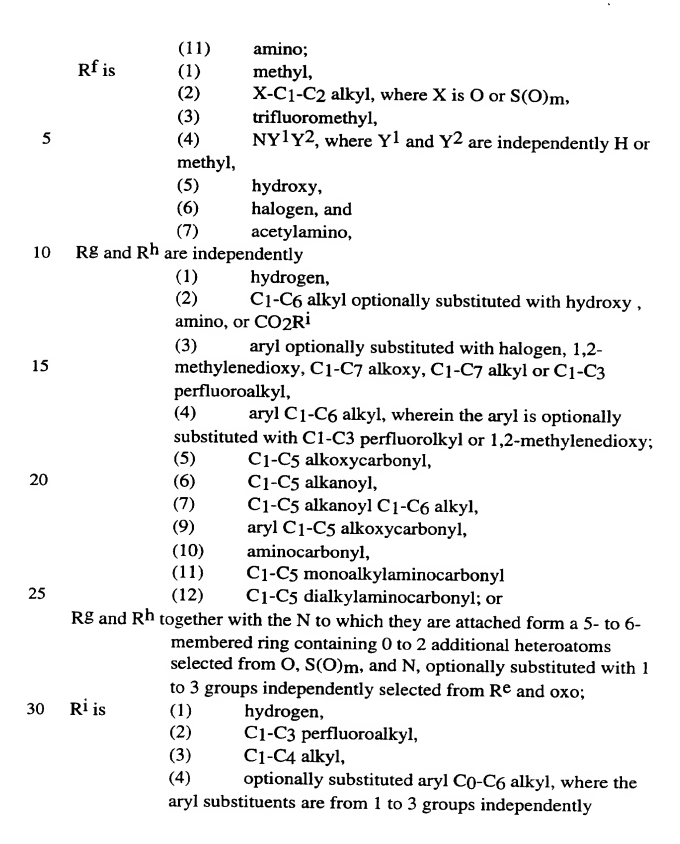
			(xx) -NRgRh,
			(xxi) 5 to 6-membered heterocycle, which may
		be satu	rated or partially unsaturated, containing from 1 to 4
		heteroa	atoms independently selected from oxygen, sulfur and
5		nitroge	en, and optionally substituted with 1 to 5 groups
		indepe	ndently selected from Re, and
			(xxii) optionally substituted aryl, wherein the
		aryl sul	bstituents are 1,2-methylenedioxy or 1 to 5 groups
		indepe	ndently selected from Re;
10	Re is	(1)	halogen,
		(2)	C ₁ -C ₃ alkyl,
			C ₁ -C ₃ perfluoroalkyl,
		(4)	$-S(O)_{\mathbf{m}}R^{\mathbf{i}},$
		(5)	cyano,
15		(6)	amino,
		(7)	RiO(CH ₂) _v -,
		(8)	RiCO ₂ (CH ₂) _v -,
		(9)	$R^{i}OCO(CH_{2})_{V},$
20		(10)	optionally substituted aryl where the substituents
20		are from	n 1 to 3 of halogen, C1-C3 alkyl, C1-C3 alkoxy, or
		hydroxy	
	nf:	(11)	SO ₂ NRgRh;
	Rf is	(1)	methyl,
25		(2)	X-C ₁ -C ₂ alkyl, where X is O or $S(O)_m$,
25		(3)	halogen,
		(4)	acetylamino,
		(5)	trifluoromethyl,
		(6)	NY^1Y^2 , where Y^1 and Y^2 are independently H or
30		methyl,	
30	Rg and Rh	(7)	hydroxy;
	KS and K"	_	•
		(1) (2)	hydrogen,
			C1-C6 alkyl optionally substituted with hydroxy,

		(3)	aryl optionally substituted with halogen, 1,2-
		methyle	enedioxy, C1-C7 alkoxy, C1-C7 alkyl or C1-C3
		perfluo	
		(4)	aryl C1-C6 alkyl, wherein the aryl is optionally
5		substitu	ted with C1-C3 perfluorolkyl or 1,2-methylenedioxy
		(5)	C1-C5 alkoxycarbonyl,
		(6)	C ₁ -C ₅ alkanoyl,
		(7)	C1-C5 alkanoyl C1-C6 alkyl,
		(9)	aryl C1-C5 alkoxycarbonyl,
10		(10)	aminocarbonyl,
		(11)	C ₁ -C ₅ monoalkylaminocarbonyl
			C ₁ -C ₅ dialkylaminocarbonyl; or
	Rg and Rh	together v	with the N to which they are attached form a 5- to 6-
		member	red ring containing 0 to 2 additional heteroatoms
15		selected	from O, S(O) _m , and N, optionally substituted with 1
	<i>-</i> .	to 3 gro	ups independently selected from Re and oxo;
	R ⁱ is	(1)	hydrogen,
		(2)	C ₁ -C ₃ perfluoroalkyl,
		(3)	C ₁ -C ₄ alkyl,
20		(4)	optionally substituted aryl C ₀ -C ₄ alkyl, where the
			stituents are from 1 to 3 groups independently
		selected	from halogen, C ₁ -C ₄ alkyl, C ₁ -C ₄ alkoxy, and
		hydroxy	
	all other va	riables are	e as defined under Formula I.
25			
			er preferred embodiment, the present invention
		_	of Formula I wherein
	R ₁ is	(1)	hydrogen,
		(2)	optionally substituted C1-C3 alkyl,
30		(3)	optionally substituted C2-C3 alkenyl,
		(4)	optionally substituted C2-C3 alkynyl,
		where th	e substitutents on the alkyl, alkenyl, and alkynyl are
		1 to 3 gr	oups independently selected from
			(i) methyl,

			(ii) X-methyl, where X is O or S(O) _m and
			(iii) halogen,
		(5)	aryl C0-C1 alkyl wherein said aryl is optionally
		substitu	ated with 1 to 3 groups independently selected from
5		Rf,	, , , , , , , , , , , , , , , , , , ,
		(6)	trifluoromethyl
	R8 is	(1)	Н,
		(2)	OH, or
		(3)	NH ₂
10	R9 is	(1)	H, or
		(2)	OH;
	R ₁₀ is	(1)	$C(O)OR^b$,
		(2)	$C(O)N(OR^b)R^c$,
		(3)	C(O)NRcRd,
15		(4)	NHC(O)ORb,
		(5)	NHC(O)NRCRd,
		(6)	CH ₂ OR ^a ,
		(7)	CH ₂ OCO ₂ Rb,
		(8)	CH ₂ OC(O)NRcRd,
20		(9)	C(O)NRCNRCRd, or
		(10)	$C(O)NR^{c}SO_{2}R^{b};$
	Ra is	(1)	hydrogen,
		(2)	optionally substituted C1-C4 alkyl,
25		(3)	optionally substituted C3-C4 alkenyl,
25		(4)	optionally substituted C3-C4 alkynyl,
		(5)	optionally substituted C1-C4 alkanoyl,
		(6)	optionally substituted aroyl,
		(7)	optionally substituted C5-C6 cycloalkanoyl,
20		(8)	optionally substituted C5-C6 cycloalkenoyl,
30		(9)	optionally substituted C1-C3 alkylsulfonyl
		where th	e substituents on the alkyl, alkenyl, alkynyl,
		alkanoy	, aroyl, cycloalkanoyl, cycloalkenoyl, and
		alkylsuli	fonyl, are from 1 to 5 groups independently selected

		from hy	droxy, C ₁	-C2 alkoxy, aryl C1-C3 alkoxy, NRgRh,			
		CO ₂ Rb,	CONRC	Rd and halogen,			
		(10)	trifluoro	omethyl,			
		(11)	arylsulf	onyl optionally substituted with 1 to 3			
5		groups i	ndepende	ently selected from methyl, trifluoromethyl			
		and halo	alogen,				
		(12)	a 5- or 6	5-membered heterocycle containing 1 to 4			
		heteroat	oms selec	ted from oxygen, sulfur and nitrogen			
		optional	ly substiti	uted by 1 to 4 groups independently			
10		selected	ed from methyl, trifluoromethyl, C(O)NRcRd, CO2Rb				
		and halo	nd halogen, and which may be saturated or partly				
	_	unsatura	saturated;				
	Rb is	(1)	Н,				
		(2)	optional	ly substituted aryl,			
15		(3)	optional	ly substituted C1-C6 alkyl,			
		(4)	optional	ly substituted C3-C6 alkenyl,			
		(5)	optional	ly substituted C3-C6 alkynyl,			
		(6)	optional	ly substituted C5-C6 cycloalkyl,			
		(7)	optional	ly substituted C5-C6 cycloalkenyl, or			
20		(8)		ly substituted 5- to 6-membered			
			ocycle containing from 1 to 4 heteroatoms				
		independently selected from oxygen, sulfur and nitrogen;					
		where the substituents on the aryl, alkyl, alkenyl, cycloalkyl,					
		cycloalkenyl, heterocycle, or alkynyl are from 1 to 10 groups					
25		independently selected from					
		(i) hydroxy,					
			(ii)	C ₁ -C ₃ alkyl,			
			(iii)	oxo,			
			(iv)	SO ₂ NRgR ^h ,			
30			(v)	aryl C ₁ -C ₃ alkoxy,			
			(vi)	hydroxy C ₁ -C ₄ alkyl,			
			(vii)	C ₁ -C ₄ alkoxy,			
			(viii)	hydroxy C ₁ -C ₄ alkoxy,			
			(ix)	amino C ₁ -C ₄ alkoxy,			

			(x)	cyano,		
			(xi)	C ₁ -C ₄ alkyl-S(O)m,		
			(xii)	C5-C6 cycloalkyl optionally substituted		
			with 1	to 4 groups independently selected from Re		
5			(xiii)	C5-C6 cycloalkenyl,		
			(xiv)	<u>▼</u> :		
			(xv)	C1-C3 alkanoyloxy,		
				C(O)NRgRh		
				CO ₂ Ri,		
10			(xvii)	-NRgRh		
			(xix)	5 to 6-membered heterocycle, which may		
		be sate	urated or pa	artially unsaturated, containing from 1 to 4		
heteroatoms independently selected from oxyg				pendently selected from oxygen, sulfur and		
		nitrog	nitrogen, and optionally substituted with 1 to 5 groups			
15		indepe	endently sel	ected from Re,		
			(xx)	optionally substituted aryl, wherein the		
		aryl su	ıbstituents a	are 1,2-methylenedioxy or 1 to 5 groups		
		indepe	endently sel	ected from Re,		
			(xxi)	optionally substituted aryl C1-C3 alkoxy,		
wherein the aryl substituents are 1,2-me 5 groups independently selected from R				ubstituents are 1,2-methylenedioxy or 1 to		
				lently selected from Re, and		
			(xxii)	C ₁ -C ₃ perfluoroalkyl;		
	Re is	(1)	halogen,			
		(2)	C1-C3 a			
25				erfluoroalkyl,		
		(4)	$-S(O)_{\mathbf{m}}F$	R ⁱ ,		
		(5)	cyano,			
		(6)	RiO(CH	2) _V -,		
• •		(7)	RiCO ₂ (C	CH ₂) _v -,		
30		(8)	RiOCO($CH_2)_{v}$,		
		(9)	optionall	y substituted aryl where the substituents		
		are from	from 1 to 3 of halogen, C ₁ -C ₃ alkyl, C ₁ -C ₃ alkoxy, or			
		hydrox	у,			
		(10)	SO2NRg	Rh or		



selected from halogen, C_1 - C_6 alkyl, C_1 - C_6 alkoxy, and hydroxy; and

all other variables are as defined under Formula I.

In another aspect of the present invention there are provided compounds having the formula X

where R₁ - R₆, R₈ and R₉ are as defined under formula I; and

10 R₁₁ is (1) COCl,

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(2) CON3, or

(3) NCO.

Compounds of formula X are useful as intermediates in the preparation of certain compounds of formula I from Compounds A, B and C.

The present invention provides in another aspect pharmaceutical compositions comprising a compound of Formula I and a pharmaceutically acceptable carrier. Such compositions may further comprise one or more other active ingredients such as anthelmintic agents, insect regulators, ecdosyne agonists and fipronil.

The present invention provides in another aspect a method for treating parasitic diseases in a mammal which comprises administering an antiparasitic amount of a compound of Formula I. The treatment may further comprise co-administering one or more other active ingredients such as anthelmintic agents, insect regulators, ecdosyne agonists and fipronil.

"Alkyl" as well as other groups having the prefix "alk", such as alkoxy, alkanoyl, alkenyl, alkynyl and the like, means carbon chains which may be linear or branched or combinations thereof. Examples of alkyl groups include methyl, ethyl, propyl, isopropyl, butyl, sec- and tertbutyl, pentyl, hexyl, heptyl and the like. "Alkenyl", "alkynyl" and other like terms include carbon chains containing at least one unsaturated C-C bond.

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The term "cycloalkyl" means carbocycles containing no heteroatoms, and includes mono-, bi- and tricyclic saturated carbocycles, as well as benzofused carbocycles. Examples of cycloalkyl include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, decahydronaphthalene, adamantane, indanyl, indenyl, fluorenyl, 1,2,3,4-tetrahydronaphalene and the like. Similarly, "cycloalkenyl" means carbocycles containing no heteroatoms and at least one non-aromatic C-C double bond, and include mono-, bi- and tricyclic partially saturated carbocycles, as well as benzofused cycloalkenes. Examples of cycloalkenyl include cyclohexenyl, indenyl, and the like.

The term "halogen" is intended to include the halogen atoms fluorine, chlorine, bromine and iodine.

The term "heterocycle", unless otherwise specfied, means 20 mono- or bicyclic compounds that are saturated or partly unsaturated, as well as benzo- or heteroaromatic ring fused saturated heterocycles or partly unsaturated heterocycles, and containing from 1 to 4 heteroatoms independently selected from oxygen, sulfur and nitrogen. Examples of saturated heterocycles include morpholine, thiomorpholine, piperidine, 25 piperazine, tetrahydropyran, tetrahydrofuran, dioxane, tetrahydrothiophene, oxazolidine, pyrrolidine; examples of partly unsaturated heterocycles include dihydropyran, dihydropyridazine, dihydrofuran, dihydrooxazole, dihydropyridine, dihydropyridazine and the like. Examples of benzo- or heteroaromatic 30 ring fused heterocycle include 2,3-dihydrobenzofuranyl, benzopyranyl, tetrahydroquinoline, tetrahydroisoquinoline, benzomorpholinyl, 1,4benzodioxanyl, 2,3-dihydrofuro(2,3-b)pyridyl and the like.

The term "aryl" is intended to include mono- and bicyclic aromatic and heteroaromatic rings containing from 0 to 5 heteroatoms independently selected from nitrogen, oxygen and sulfur. The term "aryl" is also meant to include benzofused cycloalkyl, benzofused cycloalkenyl, and benzofused heterocyclic groups. Examples of "aryl" groups include phenyl, pyrrolyl, isoxazolyl, pyrazinyl, pyridinyl, oxazolyl, thiazolyl, imidazolyl, triazolyl, tetrazolyl, furanyl, triazinyl, thienyl, pyrimidinyl, pyridazinyl, pyrazinyl, naphthyl, benzoxazolyl, benzothiazolyl, benzimidazolyl, benzofuranyl, furo(2,3-B)pyridyl, 2,3-dihydrofuro(2,3-b)pyridyl, benzoxazinyl, benzothiophenyl, quinolinyl, indolyl, 2,3-dihydrobenzofuranyl, benzopyranyl, 1,4-benzodioxanyl, indanyl, indenyl, fluorenyl, 1,2,3,4-tetrahydronaphthalene and the like.

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Aroyl means arylcarbonyl in which aryl is as defined above. Examples of NRcRd or NRgRh forming a 3- to 10-

membered ring containing 0 to 2 additional heteroatoms selected from O, S(O)_m and N are aziridine, azetidine, pyrrolidine, piperidine, thiomorpholine, morpholine, piperazine, octahydroindole, tetrahydroisoquinoline and the like.

The term "optionally substituted" is intended to include both substituted and unsubstituted; thus, for example, optionally substituted aryl could represent a pentafluorophenyl or a phenyl ring.

Certain of the above defined terms may occur more than once in the above formula and upon such occurrence each term shall be defined independently of the other; thus, for example, OR^a at C4 may represent OH and at C20 represent O-acyl.

Compounds described herein contain one or more asymmetric centers and may thus give rise to diastereomers and optical isomers. The present invention is intended to include all possible diastereomers as well as their racemic and resolved, enantiomerically pure forms and all possible geometric isomers. In addition, the present invention includes all pharmaceutically acceptable salts thereof. The term "pharmaceutically acceptable salts" refers to salts prepared from pharmaceutically acceptable non-toxic bases including inorganic bases and organic bases. Salts derived from inorganic bases include aluminum,

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ammonium, calcium, copper, ferric, ferrous, lithium, magnesium, manganic salts, manganous, potassium, sodium, zinc, and the like. Particularly preferred are the ammonium, calcium, magnesium, potassium, and sodium salts. Salts derived from pharmaceutically acceptable organic non-toxic bases include salts of primary, secondary, and tertiary amines, substituted amines including naturally occurring substituted amines, cyclic amines, and basic ion exchange resins, such as arginine, betaine, caffeine, choline, N,N--dibenzylethylenediamine, diethylamine, 2-diethylaminoethanol, 2-dimethylaminoethanol, ethanolamine, ethylenediamine, N-ethyl-morpholine, N-ethylpiperidine, glucamine, glucosamine, histidine, hydrabamine, isopropylamine, lysine, methylglucamine, morpholine, piperazine, piperidine, polyamine resins, procaine, purines, theobromine, triethylamine, trimethylamine, tripropylamine, tromethamine, and the like.

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When the compound of the present invention is basic, salts may be prepared from pharmaceutically acceptable non-toxic acids, including inorganic and organic acids. Such acids include acetic, benzenesulfonic, benzoic, camphorsulfonic, citric, ethanesulfonic, fumaric, gluconic, glutamic, hydrobromic, hydrochloric, isethionic, lactic, maleic, malic, mandelic, methanesulfonic, mucic, nitric, pamoic, pantothenic, phosphoric, succinic, sulfuric, tartaric, p-toluenesulfonic acid, and the like. Particularly preferred are citric, hydrobromic, hydrochloric, maleic, phosphoric, sulfuric, and tartaric acids.

Compounds of the present invention are named based on the trivial name of the parent compound, nodulisporic acid (compound A), and their position numbers are those as indicated in formula I.

Compounds of the present invention are prepared from the three nodulisporic acids (Compounds A, B and C), which in turn are obtained from the fermentation culture of *Nodulisporium* sp. MF-5954 (ATCC 74245). The description of the producing microorganism, the fermentation process, and the isolation and purification of the three nodulisporic acids are disclosed in US Patent 5,399,582, issued March 21, 1995, which is hereby incorporated by reference in its entirety.

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The above structural formula is shown without a definitive stereochemistry at certain positions. However, during the the course of the synthetic procedures used to prepare such compounds, or using racemization or epimerization procedures known to those skilled in the art, the products of such procedures can be a mixture of stereoisomers. In particular, the stereoisomers at C1, C4, C20, C26, C31 and C32 may be oriented in either the alpha- or beta-position, representing such groups oriented below or above the plane of the molecule, respectively. In each such case, and at other positions in the molecule, both the alpha- and beta-configurations are intended to be included within the ambit of this invention.

Compounds of formula I wherein the allyl group at position 26 is in the epi configuration may be obtained by treatment of the appropriate precursor with a bases such as hydroxide, methoxide, imidazole, triethylamine, potassium hydride, lithium diisopropylamide and the like in protic or aprotic solvents (as appropriate) such as water, methanol, ethanol, methylene chloride, chloroform, tetrahydrofuran, dimethylformamide and the like. The reaction is complete at temperatures from -78°C to the reflux temperature of the solution in from 15 minutes to 12 hours.

Compounds of formula I where R2 (and R1 is hydrogen), R3, R4 and R8 independently are hydroxy may be inverted by treatment of the appropriate alcohol using protocols known to those skilled in the art. For example, the alcohol may be reacted under Mitsunobu conditions with a carboxylic acid (formic acid, propionic acid, 2-chloroacetic acid, 25 benzoic acid, para-nitro-benzoic acid and the like), a tri-substituted phosphine (triphenylphosphine, tri-n-butylphoshine, tripropylphosphine and the like) and a dialkyl diazodicarboxylate (diethyl diazodicarboxylate, dimethyl diazodicarboxylate, diisopropyl diazodicarboxylate and the like) in an aprotic solvent such as methylene 30 chloride, tetrahydrofuran, chloroform, benzene and the like. The Mitsunobu reactions are complete in from 1 to 24 hours at temperatures from 0°C to the reflux temperature of the solution. The resultant esters may be hydrolyzed by treatment with hydroxide or ammonium hydroxide

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in a protic solvent such as methanol, ethanol, water, tetrahydrofuran/water or dimethylformamide/water and the like at from 0°C to the reflux temperature of the solution. Alternatively, the resultant esters may be hydrolyzed by treatment with a Lewis acid, such as magnesium chloride, aluminum chloride, titanium tetra-isopropoxide and the like in a protic solvent such as methanol, ethanol, isopropanol and the like and the reactions are complete in from 1 to 24 hours at 0°C to the reflux temperature of the solution.

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During certain reactions described below, it may be necessary to protect the groups at R2, R3, R4, R8, R9 and R10. With these positions protected, the reactions may be carried out at other positions without affecting the remainder of the molecule. Subsequent to any of the described reactions (vida infra), the protecting group(s) may be removed and the unprotected product isolated. The protecting groups employed at R2, R3, R4, R8, R9 and R10 are those which may be readily synthesized, not significantly affected by the reactions at the other positions, and may be removed without significantly affecting any other functionality of the molecule. One preferred type of protecting group is the tri-substituted silyl group, preferably the tri-loweralkyl silyl group or di-loweralkyl-aryl silyl group. Especially preferred examples are the trimethylsilyl, triethylsilyl, triisopropylsilyl, tert-butyldimethylsilyl and dimethylphenylsilyl groups.

The protected compound may be prepared with the appropriately substituted silyl trifluoromethanesulfonate or silyl halide, preferably the silyl chloride. The reaction is carried out in an aprotic solvent such as methylene chloride, benzene, toluene, ethyl acetate, isopropyl acetate, tetrahydrofuran, dimethylformamide and the like. In order to minimize side reactions, there is included in the reaction mixture a base to react with the acid released during the course of the reaction. Preferred bases are amines such as imidazole, pyridine, triethylamine or diisopropylethylamine and the like. The base is required in amounts equimolar to the amount of hydrogen halide liberated, however, generally several equivalents of the amine are employed. The reaction is stirred at

from 0°C to the reflux temperature of the reaction mixture and is complete from 1 to 24 hours.

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The silyl group is removed by treatment of the silyl compound with anhydrous pyridine-hydrogen fluoride in tetrahydrofuran or dimethylsulfoxide or with tetraalkylammonium fluoride in tetrahydrofuran. The reaction is complete in from 1 to 24 hours at from 0°C to 50°C. Alternatively, the silyl group may be removed by stirring the silylated compound in lower protic solvents such as methanol, ethanol, isopropanol and the like catalyzed by an acid, preferably a sulfonic acid monohydrate such as <u>para</u>-toluenesulfonic acid, benzenesulfonic acid or carboxylic acids such as acetic acid, propionic acid, monochloroacetic acid, dichloroacetic acid, trichloroacetic acid and the like. The reaction is complete in 1 to 24 hours at from 0°C to 50°C.

Protecting groups that may also be suitably used in the preparation of compounds of the present invention may be found in standard textbooks such as Greene and Wutz, <u>Protective Groups in Organic Synthesis</u>, 1991, John Wiley & Sons, Inc.

Compounds of formula I where R₁ and R₂ together represent an oxime, =NORa, may be prepared by treating the appropriate oxo analog with H₂NORa to produce the corresponding oxime. Oxime formation may be accomplished using techniques known to those skilled in the art, including, but not restricted to, the use of H₂NORa either as the free base or as an acid addition salt such as the HCl salt, or an O-protected hydroxylamine such as O-trialkylsilylhydroxylamine, in a protic solvent such as methanol, ethanol, isopropanol and the like or aprotic solvents such as methylene chloride, chloroform, ethyl acetate, isopropyl acetate, tetrahydrofuran, dimethylformamide, benzene, toluene and the like, as appropriate. The reactions may by catalyzed by the addition of sulfonic acids, carboxylic acids or Lewis acids, including, but not limited to, benzenesulfonic acid monhydrate, para-toluenesulfonic acid monohydrate, acetic acid, zinc chloride and the like.

Similarly, compounds of formula I wherein R₁ and R₂ together represent =NNR^cR^d may be prepared by treating the appropriate

oxo analog with H2NNR^cR^d to give the corresponding hydrazones using conditions directly analogous to those described for oxime formation.

Compounds of formula I wherein one or both of the ____ bonds represent a single bond may be prepared from the corresponding compound wherein ____ is a double bond by conventional hydrogenation procedures. The double bonds may be hydrogenated with any of a variety of standard precious metal hydrogenation catalysts such as Wilkinson's catalyst, Pearlman's catalyst, 1-25% palladium on carbon, 1-25% platinum on carbon and the like. The reaction is generally carried out in a non-reducible solvents (either protic or aprotic) such as methanol, ethanol, isopropanol, tetrahydrofuran, ethyl acetate, isopropyl acetate, benzene, toluene, dimethylformamide and the like. The hydrogen source may be hydrogen gas from 1 to 50 atmospheres of pressure or other hydrogen sources such as ammonium formate, cyclohexene, cyclohexadiene and the like. The reduction also may be carried out using sodium dithionite and sodium bicarbonate in the presence of a phase transfer catalyst, in particular a tetraalkylammonium phase transfer catalyst, and the like. The reactions may be run from 0°C to 100°C and are complete in from 5 min to 24 hours.

Compounds of formula I wherein R8 and R9 are both hydroxyl groups may be prepared according to the procedure shown in Scheme I.

SCHEME I

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Thus, Compound II is treated with osmium tetroxide under conditions known to those skilled in the art to yield the diol product III. Also produced during this reaction is the aldehyde IV. Osmium tetroxide may

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be used either stoichiometrically or catalytically in the presence of an oxidant, including, but not restricted to, morpholine N-oxide, trimethylamine N-oxide, hydrogen peroxide, tert-butyl hydroperoxide and the like. The dihydroxylation reactions may be performed in a variety of solvents or mixtures of solvents. These include both protic and aprotic solvents such as water, methanol, ethanol, tert-butanol, ether, tetrahydrofuran, benzene, pyridine, acetone and the like. The reactions may be performed at from -78°C to 80°C and are complete in from 5 minutes to 24 hours.

Compounds of formula I wherein R8 is NRcRd and R9 is hydrogen may be prepared by treatment of the appropriate precursor containing the C31-C32 unsaturation with HNRcRd or HCl• HNRcRd in an appropriate protic or aprotic solvents such as methanol, ethanol, benzene, toluene, dimethylformamide, dioxane, water and the like. The reaction may be facilitated by the addition of bases such as pyridine, triethylamine, sodium carbonate and the like or Lewis acids such as zinc chloride, magnesium chloride and the like. The reactions are complete in from 1 to 24 hours at temperatures from 0°C to the reflux temperature of the solution.

20 Compounds of formula I wherein R2 is OH and R1 is H may be prepared from the corresponding ketone by treating the appropriate oxo analog with standard reducing agents including, but not restricted to, sodium borohydride, lithium borohydride, lithium aluminum hydride, potassium tri-sec-butyl borohydride, diisobutylaluminum hydride, diborane oxazaborolidines and alkylboranes (both achiral and chiral). 25 These reactions are performed in a manner known to those skilled in the art and are carried out in non-reducible solvents such as methanol, ethanol, diethyl ether, tetrahydrofuran, hexanes, pentane, methylene chloride and the like. The reactions are complete in from 5 minutes to 24 hours at temperatures ranging from -78°C to 60°C. Compounds of 30 formula I wherein R2 is OH, R1 is H and R10 is CH2OH may be obtained by reacting the appropriate carboxylic acid or ester analog (e.g., where R₁₀ is CO₂H or CO₂R^a) with the more reactive reducing agents as described above, including lithium aluminum hydride, lithium

borohydride and the like. Compounds of formula I wherein R₂ and R₁ together are oxo and R₁₀ is CH₂OH may be obtained by reacting the appropriate carboxylic acid (e.g., where R₁₀ is CO₂H) with less reactive reducing agents such as diborane and the like.

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Compounds of formula I wherein R₂ is OH and R₁ is other than H, may be prepared from the corresponding ketone by treating the appropriate oxo analog with a Grignard reagent R₁MgBr, or with a lithium reagent R₁Li. These reactions are performed in a manner known to those skilled in the art and preferably are performed in aprotic solvents such as diethyl ether, tetrahydrofuran, hexanes or pentanes. The reactions are complete in from 5 minutes to 24 hours at temperatures ranging from -78°C to 60°C.

Compounds of formula I where R₁₀ is C(O)N(OR^b)R^c or C(O)NR^cR^d are prepared from the corresponding carboxylic acid using standard amide-forming reagents known to those skilled in the art. The reaction is carried out using at least one equivalent of an amine nucleophile, HN(OR^b)R^c or HNR^cR^d, although preferably ten to one hundred equivalents of amine nucleophiles are employed. Amideforming reagents include, but are not restricted to,

- dicyclohexylcarbodiimide, 1-(3-dimethylaminopropyl)-3ethylcarbodiimide hydrochloride (EDC•HCl), diisopropylcarbodiimide, benzotriazol-1-yloxy-tris(dimethylamino)phosphonium hexafluorphosphate (BOP), bis(2-oxo-3-oxazolidinyl)phosphinic chloride (BOP-Cl), benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium
- hexafluorophosphate (PyBOP), chloro-tris-pyrrolidino-phosphonium hexafluorophosphate (PyCloP), bromo-tris-pyrrolidino-phosphonium hexafluorophosphate (PyBroP), diphenylphosphoryl azide (DPPA), 2-(1H-benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU), O-benzotriazol-1-yl-N,N,N',N'-bis(pentamethylene)uronium
- hexafluorophosphate and 2-chloro-1-methylpyridinium iodide. The amide-forming reactions may be facilitated by the optional addition of N-hydroxybenzotriazole or N-hydroxy-7-aza-benzotriazole. The amidation reaction is generally performed using at least one equivalent (although several equivalents may be employed) of amine bases such as

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triethylamine, diisopropylethylamine, pyridine, N,N-dimethylaminopyridine and the like. The carboxyl group may be activated for amide bond formation via its corresponding acid chloride or mixed anhydride, using conditions known to those skilled in the art. These amide-forming reactions are carried out in aprotic solvents such as

These amide-forming reactions are carried out in aprotic solvents such as methylene chloride, tetrahydrofuran, diethyl ether, dimethylformamide, N-methylpyrrolidine and the like at -20°C to 60°C and are complete in 15 minutes to 24 hours.

Compounds of formula I where R₁₀ is cyano may be

prepared by treatment of the appropriate carboxamide with dehydrating reagents known to those skilled in the art such as <u>para</u>-toluenesulfonyl chloride, methanesulfonyl chloride, acetyl chloride, thionyl chloride, phosphorus oxychloride or catecholboron chloride in an aprotic solvent such as methylene chloride, chloroform, tetrahydrofuran, benzene,

toluene and the like. The reactions are complete in from 15 minutes to 24 hours at temperatures from -78°C to the reflux temperature of the solution.

Compounds of formula I where R₁₀ is C(O)OR^b are prepared from the corresponding carboxylic acid using standard esterforming reagents known to those skilled in the art. The esterification reaction is carried out using at least one equivalent of an alcohol, HOR^b, although preferably ten to one hundred equivalents of alcohol are used; the esterification also may be carried out using the alcohol as solvent. Esterification reagents include, but are not restricted to,

- dicyclohexylcarbodiimide, 1-(3-dimethylaminopropyl)-3ethylcarbodiimide hydrochloride (EDC•HCl), diisopropylcarbodiimide, benzotriazol-1-yloxy-tris(dimethylamino)phosphonium hexafluorphosphate (BOP), bis(2-oxo-3-oxazolidinyl)phosphinic chloride (BOP-Cl), benzotriazole-1-yl-oxy-tris-pyrrolidino-phosphonium
- hexafluorophosphate (PyBOP), chloro-tris-pyrrolidino-phosphonium hexafluorophosphate (PyCloP), bromo-tris-pyrrolidino-phosphonium hexafluorophosphate (PyBroP), diphenylphosphoryl azide (DPPA), 2-(1H-benzotriazole-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU), O-benzotriazol-1-yl-N,N,N',N'-bis(pentamethylene)uronium

hexafluorophosphate and 2-chloro-1-methylpyridinium iodide. The ester-forming reactions may be facilitated by the optional addition of N-hydroxybenzotriazole, N-hydroxy-7-aza-benzotriazole, 4-(N,N-dimethylamino)pyridine or 4-pyrrolidinopyridine. The reaction is generally performed using at least one equivalent (although several equivalents may be employed) of amine bases such as triethylamine, diisopropylethylamine, pyridine and the like. The carboxyl group may be activated for ester bond formation via its corresponding acid chloride or mixed anhydride, using conditions known to those skilled in the art.

These ester-forming reactions are carried out in aprotic solvents such as methylene chloride, tetrahydrofuran, diethyl ether, dimethylformamide, N-methylpyrrolidine and the like at temperatures ranging from -20°C to

60°C and are complete in 15 minutes to 24 hours.

Compounds of formula I wherein one or more of R2, R3, R4, R8 and R9 is ORa, OCO2Rb or OC(O)NRcRd, and/or where R10 is 15 CH2ORa, CH2OCO2Rb or CH2OC(O)NRcRd may be prepared using known methods for acylation, sulfonylation and alkylation of alcohols. Thus, acylation may be accomplished using reagents such as acid anhydrides, acid chlorides, chloroformates, carbamoyl chlorides, isocyanates and amine bases according to general procedures known to 20 those skilled in the art. Sulfonylations may be carried out using sulfonylchlorides or sulfonic anhydrides. The acylation and sulfonylation reactions may be carried out in aprotic solvents such as methylene chloride, chloroform, pyridine, benzene, toluene and the like. The acylation and sulfonylation reactions are complete in from 15 minutes to 25 24 hours at temperatures ranging from -20°C to 80°C. The degree of acylation, sulfonylation and alkylation will depend on the amount of the reagents used. Thus, for example, using one equivalent of an acylating reagent and one equivalent of nodulisporic acid results in a product mixture containg 4- and 20-acylated nodulisporic acid; such a mixture 30 may be separated by conventional techniques such as chromatography.

Compounds of formula I wherein one or more of R2, R3, R4, R8 and R9 is OR^a and/or where R10 is CH2OR^a, may be prepared using methods known to those skilled in the art for the alkylation of

alcohols. Thus, alkylation may be accomplished using reagents including, but not restricted to, halides IRa, BrRa, ClRa, diazo reagents N2Ra, trichloroacetimidates RaOC(NH)CCl3, sulfates RaOSO2Me, RaOSO2CF3, and the like. The alkylation reactions may be facilitated by the addition of acid, base or Lewis acids, as appropriate. The reactions are performed in aprotic solvents such as methylene chloride, chloroform, tetrahydrofuran, benzene, toluene, dimethylformamide, N-methylpyrrolidine, dimethyl sulfoxide, hexamethylphosphoramide and are complete at from 0°C to the reflux temperature of the solution from 15 minutes to 48 hours.

Compounds of formula I where R₁₀ is NHC(O)OR^b or C(O)NR^cR^d are prepared from the corresponding carboxylic acid via the corresponding acyl azide (VI) and isocyanate (VII) as shown in Scheme II.

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SCHEME II

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In Scheme II, R8, R9, Rb, Rc, Rd and — have the same meaning as defined under formula I. Thus, the carboxylic acid (compound V) is treated with diphenylphosphoryl azide to provide the acyl azide (compound VI). Heating of compound VI in an aprotic solvent such as benzene, toluene, dimethylformamide and the like results in a rearrangement yielding compound VII, an isocyanate. Compound VII may be reacted in an aprotic solvent such as benzene, toluene, methylene chloride, 1,2-dichloroethylene, dimethylformamide and the like, with an alcohol RbOH, such as methanol, ethanol, benzyl alcohol, 2-trimethylsilylethanol, 2,2,2-trichloroethanol, methyl glyocolate, phenol and the like to yield compound VIII, a carbamate. The addition of one or

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more equivalents of an amine base such as triethylamine, diisopropylethylamine, pyridine and the like may be employed to accelerate carbamate formation. The carbamate-forming reactions may be performed from 0°C to 100°C and are complete in 15 minutes to 24 hours.

Compounds of formula IX may be prepared when compounds of formula VII are reacted with an appropriate amine HNRcRd in an aprotic solvent such as methylene chloride, tetrahydrofuran, dimethylformamide, dimethylsulfoxide, benzene, toluene and the like. The urea-forming reactions may be performed from 0°C to 100°C and are complete in 15 minutes to 24 hours.

The instant compounds are potent endo- and ectoantiparasitic agents against parasites particularly helminths, ectoparasites, insects, and acarides, infecting man, animals and plants, thus having utility in human and animal health, agriculture and pest control in household and commercial areas.

The disease or group of diseases described generally as helminthiasis is due to infection of an animal host with parasitic worms known as helminths. Helminthiasis is a prevalent and serious economic problem in domesticated animals such as swine, sheep, horses, cattle, goats, dogs, cats, fish, buffalo, camels, llamas, reindeer, laboratory animals, furbearing animals, zoo animals and exotic species and poultry. Among the helminths, the group of worms described as nematodes causes widespread and often times serious infection in various species of animals. The most common genera of nematodes infecting the animals referred to above are Haemonchus, Trichostrongylus, Ostertagia, Nematodirus, Cooperia, Ascaris, Bunostomum, Oesophagostomum, Chabertia, Trichuris, Strongylus, Trichonema, Dictyocaulus, Capillaria, Habronema, Druschia, Heterakis, Toxocara, Ascaridia, Oxyuris, Ancylostoma, Uncinaria, Toxascaris and Parascaris. Certain of these, such as Nematodirus, Cooperia, and Oesophagostomum attack primarily the intestinal tract while others, such as Haemonchus and Ostertagia, are

more prevalent in the stomach while still others such as Dictyocaulus are found in the lungs. Still other parasites may be located in other tissues

and organs of the body such as the heart and blood vessels, subcutaneous and lymphatic tissue and the like. The parasitic infections known as helminthiases lead to anemia, malnutrition, weakness, weight loss, severe damage to the walls of the intestinal tract and other tissues and organs and, if left untreated, may result in death of the infected host. The compounds of this invention have activity against these parasites, and in addition are also active against Dirofilaria in dogs and cats, Nematospiroides, Syphacia, Aspiculuris in rodents, arthropod ectoparasites of animals and birds such as ticks, mites such as scabies lice, fleas, blowflies, and other biting insects in domesticated animals and poultry, such as Tenophalides, Ixodes, Psoroptes, and Hemotobia, in sheep Lucilia sp., biting insects and such migrating dipterous larvae as Hypoderma sp. in cattle, Gastrophilus in horses, and Cuterebra sp. in rodents and nuisance flies including blood feeding flies and filth flies.

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The instant compounds are also useful against parasites which infect humans. The most common genera of parasites of the gastro-intestinal tract of man are Ancylostoma, Necator, Ascaris, Strongyloides, Trichinella, Capillaria, Trichuris, and Enterobius. Other medically important genera of parasites which are found in the blood or other tissues and organs outside the gastrointestinal tract are the filiarial worms such as Wuchereria, Brugia, Onchocerca and Loa, Dracunuculus and extra intestinal stages of the intestinal worms Strongyloides and Trichinella. The compounds are also of value against arthropods parasitizing man, biting insects and other dipterous pests causing annoyance to man.

The compounds are also active against household pests such as the cockroach, Blatella sp., clothes moth, Tineola sp., carpet beetle, Attagenus sp., the housefly Musca domestica as well as fleas, house dust mites, termites and ants.

The compounds are also useful against insect pests of stored grains such as Tribolium sp., Tenebrio sp. and of agricultural plants such as aphids, (Acyrthiosiphon sp.); against migratory orthopterans such as locusts and immature stages of insects living on plant tissue. The compounds are useful as a nematocide for the control of soil nematodes

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and plant parasites such as Meloidogyne sp. which may be of importance in agriculture. The compounds are also highly useful in treating acerage infested with fire ant nests. The compounds are scattered above the infested area in low levels in bait formulations which are brought back to the nest. In addition to a direct-but-slow onset toxic effect on the fire ants, the compound has a long-term effect on the nest by sterilizing the queen which effectively destroys the nest.

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The compounds of this invention may be administered in formulations wherein the active compound is intimately admixed with one or more inert ingredients and optionally including one or more additional active ingredients. The compounds may be used in any composition known to those skilled in the art for administration to humans and animals, for application to plants and for premise and area application to control household pests in either a residential or commercial setting. For application to humans and animals to control internal and external parasites, oral formulations, in solid or liquid or parenteral liquid, implant or depot injection forms may be used. For topical application dip, spray, powder, dust, pour-on, spot-on, jetting fluid, shampoos, collar, tag or harness, may be used. For agricultural premise or area application, liquid spray, powders, dust, or bait forms may be used. In addition "feed-through" forms may be used to control nuisance flies that feed or breed in animal waste. The compounds are formulated, such as by encapsulation, to lease a residue of active agent in the animal waste which controls filth flies or other arthropod pests.

These compounds may be administered orally in a unit dosage form such as a capsule, bolus or tablet, or as a liquid drench where used as an anthelmintic in mammals. The drench is normally a solution, suspension or dispersion of the active ingredient usually in water together with a suspending agent such as bentonite and a wetting agent or like excipient. Generally, the drenches also contain an antifoaming agent. Drench formulations generally contain from about 0.001 to 0.5% by weight of the active compound. Preferred drench formulations may contain from 0.01 to 0.1% by weight. The capsules and

boluses comprise the active ingredient admixed with a carrier vehicle such as starch, talc, magnesium stearate, or di-calcium phosphate.

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Where it is desired to administer the instant compounds in a dry, solid unit dosage form, capsules, boluses or tablets containing the desired amount of active compound usually are employed. These dosage forms are prepared by intimately and uniformly mixing the active ingredient with suitable finely divided diluents, fillers, disintegrating agents, and/or binders such as starch, lactose, talc, magnesium stearate, vegetable gums and the like. Such unit dosage formulations may be varied widely with respect to their total weight and content of the antiparasitic agent depending upon factors such as the type of host animal to be treated, the severity and type of infection and the weight of the host.

When the active compound is to be administered via an animal feedstuff, it is intimately dispersed in the feed or used as a top dressing or in the form of pellets or liquid which may then be added to the finished feed or optionally fed separately. Alternatively, feed based individual dosage forms may be used such as a chewable treat. Alternatively, the antiparasitic compounds of this invention may be administered to animals parenterally, for example, by intraruminal, intramuscular, intravascular, intratracheal, or subcutaneous injection in which the active ingredient is dissolved or dispersed in a liquid carrier vehicle. For parenteral administration, the active material is suitably admixed with an acceptable vehicle, preferably of the vegetable oil variety such as peanut oil, cotton seed oil and the like. Other parenteral vehicles such as organic preparation using solketal, glycerol formal, propylene glycol, and aqueous parenteral formulations are also used. The active compound or compounds are dissolved or suspended in the parenteral formulation for administration; such formulations generally contain from 0.0005 to 5% by weight of the active compound.

The agents of this invention can be used in the treatment and/or prevention of diseases caused by parasites, for example, arthropod parasites such as ticks, lice, fleas, mites and other biting arthropods in domesticated animals and poultry. The agents of this invention also are useful in the prevention and treatment of diseases caused by

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helminthiasis. They are also effective in treatment of parasitic diseases that occur in other animals including humans. The optimum amount to be employed for best results will, of course, depend upon the particular compound employed, the species of animal to be treated and the type and severity of parasitic infection or infestation. Generally good results are obtained with our novel compounds by the oral administration of from about 0.001 to 500 mg per kg of animal body weight, such total dose being given at one time or in divided doses over a relatively short period of time such as 1-5 days. With the preferred compounds of the invention, excellent control of such parasites is obtained in animals by administering from about 0.025 to 100 mg per kg of body weight in a single dose. Repeat treatments are given as required to combat re-infections and are dependent upon the species of parasite and the husbandry techniques being employed. Repeat treatments may be given daily, weekly, biweekly or monthly, or any combination thereof, as required. The techniques for administering these materials to animals are known to those skilled in the veterinary field.

When the compounds described herein are administered as a component of the feed of the animals, or dissolved or suspended in the drinking water, compositions are provided in which the active compound or compounds are intimately dispersed in an inert carrier or diluent. By inert carrier is meant one that will not react with the antiparasitic agent and one that may be administered safely to animals. Preferably, a carrier for feed administration is one that is, or may be, an ingredient of the animal ration.

Suitable compositions include feed premixes or supplements in which the active ingredient is present in relatively large amounts and which are suitable for direct feeding to the animal or for addition to the feed either directly or after an intermediate dilution or blending step. Typical carriers or diluents suitable for such compositions include, for example, distillers' dried grains, corn meal, citrus meal, fermentation residues, ground oyster shells, wheat shorts, molasses solubles, corn cob meal, edible bean mill feed, soya grits, crushed limestone and the like. The active compounds are intimately dispersed throughout the carrier by

methods such as grinding, stirring, milling or tumbling. Compositions containing from about 0.005 to 2.0% weight of the active compound are particularly suitable as feed premixes. Feed supplements, which are fed directly to the animal, contain from about 0.0002 to 0.3% by weight of the active compounds.

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Such supplements are added to the animal feed in an amount to give the finished feed the concentration of active compound desired for the treatment and control of parasitic diseases. Although the desired concentration of active compound will vary depending upon the factors previously mentioned as well as upon the particular compound employed, the compounds of this invention are usually fed at concentrations of between 0.00001 to 0.002% in the feed in order to achieve the desired anti-parasitic result.

In using the compounds of this invention, the individual compounds may be prepared and used in that form. Alternatively, mixtures of the individual compounds may be used, or they may be combined with other active compounds not related to the compounds of this invention.

The compounds of this invention are also useful in combatting agricultural pests that inflict damage upon crops while they are growing or while in storage. The compounds are applied using known techniques as sprays, dusts, emulsions and the like, to the growing or stored crops to effect protection from such agricultural pests.

Compounds of this invention may be co-administered with anthelmintic agents. These anthelmintic agents are meant to include, but not be restricted to, compounds selected from the avermectin and milbemycin class of compounds such as ivermectin, avermectin, abamectin, emamectin, eprinamectin, doramectin, fulladectin, moxidectin, Interceptor and nemadectin. Additional anthelmintic agents include the benzimidazoles such as thiabendazole, cambendazole, parbendazole, oxibendazole, mebendazole, flubendazole, fenbendazole, oxfendazole, albendazole, cyclobendazole, febantel, thiophanate and the like. Additional anthelmintic agents include imidazothiazoles and

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tetrahydropyrimidines such as tetramisole-levamisole, butamisole, pyrantel, pamoate, aoxantel or morantel.

Compounds of this invention may be co-administered with fipronil.

Compounds of this invention may be co-administered with an insect growth regulator with molt inhibiting activity such as lufenuron and the like.

Compounds of this invention may be co-administered with ecdysone agonist such as tebufenozide and the like, which induces premature molt and causes feeding to cease.

The co-administered compounds are given via routes, and in doses, that are customarily used for those compounds.

Also included in the present invention are pharmaceutical compositions containing a compound of the present invention in combination with an anthelmintic agent, fipronil, an insect growth regulator, or a ecdysone agonist.

The following examples are provided to more fully illustrate the present invention, and shall not be construed as limiting the scope in any manner.

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EXAMPLE 1 Methyl nodulisporate

To 5.4 mg nodulisporic acid in 5 mL methanol at room temperature was added 0.5 mL 10% trimethylsilyldiazomethane in hexanes. After 15 minutes, three drops of glacial acetic acid was added and the solution diluted with benzene, frozen and lyophilized. Pure methyl ester was obtained following reversed-phase HPLC purification using 85:15 methanol:water as eluant and the product was characterized by ¹H NMR and mass spectrometry.

EXAMPLE 2 Methyl 29,30-dihydro-20,30-oxa-nodulisporate

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To 0.8 mg Compound B in 1 mL methanol at room temperature was added 0.2 mL 1 M trimethylsilyldiazomethane in hexanes. After 5 minutes, 0.1 mL glacial acetic acid was added, the solution stirred for three minutes and the 2 mL saturated NaHCO₃ was added (foaming occurred). The solution was extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using 15:85 water/methanol as eluant and the purified product was characterized by ¹H NMR.

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EXAMPLE 3

Methyl 31-hydroxy-20,30-oxa-29,30,31,32-tetrahydronodulisporate

To 1 mg Compound C in 1 mL methanol at room temperature was added 0.2 mL 1 M trimethylsilyldiazomethane in hexanes. After 5 minutes, 0.1 mL glacial acetic acid was added, the solution stirred for three minutes and the 2 mL saturated NaHCO₃ was added (foaming occurred). The solution was extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using 17.5:82.5 water/methanol as eluant and the purified product was characterized by ¹H NMR.

EXAMPLE 4 Ethyl nodulisporate

To a solution containing 20 mg nodulisporic acid in 2 mL methylene chloride at room temperature was added 0.11 mL ethanol, 0.008 mL diisopropylethylamine, 1 mg N,N-dimethylaminopyridine (DMAP) followed by 13 mg BOP reagent. After 50 hours at room temperature, the solution was poured into 1/1 saturated sodium bicarbonate/brine and extracted with methylene chloride. The combined organic layers were dried over sodium sulfate, the solids were removed by filtration and the solution concentrated under reduced pressure. Pure product was obtained following preparative TLC on silica gel (one 1000 micron plate) using 1/3 acetone/hexanes as eluant. Purified product (15

mg) was characterized by proton NMR and mass spectrometry (m/z: 708.4 (M+1)).

The general procedure of Example 4 was repeated using the alcohols listed in Table 1 below to provide the corresponding nodulisporate derivatives. These compounds were characterized by proton NMR and/or mass spectrometry (m/z is for (M+1) unless otherwise specified).

10 Table 1: Ester Derivatives of Nodulisporic Acid

Ex.	m/z	Alcohol	Rb
5	797.6	N-Hydroxybenzotriazole	NN N
6	724.4	2-Hydroxyethanol	CH2CH2OH
7	807.5	2-(Diisopropylamino)- ethanol	CH2CH2N(CH(CH3)2)2
8	738.4	3-Hydroxypropanol	CH ₂ CH ₂ CH ₂ OH
9	752.3	4-Hydroxybutanol	CH2CH2CH2CH2OH
10	767.0	5-Hydroxypentanol	CH2CH2CH2CH2CH2OH
11	751.5	2-Dimethylaminoethanol	CH ₂ CH ₂ N(CH ₃) ₂
12	837.7	3-Diisopropylamino-2-	CH ₂ CH(OH)CH ₂ N(CH(CH ₃)
		hydroxypropanol	2)2
13	768.9	2-(2-Hydroxyethoxy)- ethanol	CH2CH2OCH2CH2OH

			
14	815.4	4-Nitrobenzyl alcohol	CH ₂ Ph(4-NO ₂)
15	815.4	3-Nitrobenzyl alcohol	CH ₂ Ph(3-NO ₂)
16	807.7	2-Hydroxy-3-(1- pyrrolidinyl)propanol	CH ₂ CH(OH)CH ₂ -N
17	793.7	4-(2-Hydroxyethyl)- morpholine	CH ₂ CH ₂ — N O
18	762.4	2,2,2-Trifluoroethanol	CH ₂ CF ₃
19		2-(Hydroxymethyl)furan	CH ₂ O
20	764.5	5-Hydroxypentan-2-one	CH2CH2CH2C(=O)CH3
21		3-Phenylpropanol	CH2CH2CH2Ph
22	764.3	3,3-Dimethylbutanol	CH ₂ CH ₂ C(CH ₃) ₂ CH ₃
23		2-(N-Acetylamino)-3- hydroxypyridine	NHC(O)CH ₃
24	766.7	3,4-Dihydroxytetrahydro- furan, Isomer A	OH, isomer A
25	766.6	3,4-Dihydroxytetrahydro- furan, Isomer B	OH, isomer B
26	831.5	1,1,1,3,3,3-hexafluoro- isopropanol	CH(CF3)2
27		2-(Trifluoromethyl)benzyl alcohol	CH ₂ Ph(2-CF ₃)

EXAMPLE 28
General Procedure for the Preparation of Additional Ester Derivatives of Compounds A, B and C

To a solution containing 20 mg Compound A, B or C in 2 mL methylene chloride at room temperature add 110 mg of an alcohol selected from Table 2, 0.008 mL diisopropylethylamine and 1 mg DMAP followed by 13 mg BOP reagent. After from 1 hour to 3 days at room 5 temperature, pour the solution into 1/1 saturated sodium bicarbonate/brine and extract with methylene chloride. The combined organic layers may be dried over sodium sulfate and the solids may be removed by filtration. Concentrate the solution under reduced pressure. Pure product may be obtained following flash chromatography or preparative TLC on silica gel or reversed-phase liquid chromatography. Purified product may be characterized by proton NMR and/or mass spectrometry.

- Table 2: Alcohols for the Preparation of Additional Ester Derivatives of 15 Compounds A, B and C
 - 3-(Methylthio)propanol, 1H,1H-Pentafluoropropanol, 2-Pentyn-1-ol, 3-Pentyne-1-ol, 4-Pentyne-1-ol, Propanol, 2-Hydroxyethanol, Methyl
- glycolate, Glycolic acid, 4-(Methoxy)benzyl alcohol, 3-20 (Dimethylamino)propanol, 3-(4-Morpholinyl)propanol, 2-(Hydroxymethyl)pyridine, 1-(2-Hydroxyethyl)piperazine, 2-Hydroxy-3phenylpropanol, 2-(Hydroxyethoxy)ethanol, 4-(2-Hydroxyethyl)morpholine, 1-(2-Hydroxyethyl)piperidine, 3-
- (Hydroxymethyl)pyridine, 1-(Hydroxymethyl)pyrimidine, 3-25 Hydroxypropanol, 4-Hydroxybutanol, 1-(2-Hydroxyethyl)-4methylpiperazine, 2-(2-Hydroxyethyl)pyridine, 1-(3-Hydroxypropyl)-2pyrrolidinone, 1-(2-Hydroxyethyl)pyrrolidine, 1-(3-Hydroxypropyl)imidazole, 2-Hydroxybutanol, 4-
- (Hydroxymethyl)pyridine, 2-Hydroxypyrazine, Hydroxyacetonitrile, 6-30 Hydroxyhexanol, 4-(3-Hydroxypropyl)morpholine, 2-Hydroxypropanol, 2-Hydroxypentanol, 1-Hydroxy-1-(hydroxymethyl)cyclopentane, 2-(Methylthio)ethanol, 3-Hydroxy-1,2,4-triazine, 2-Amino-3hydroxypyridine, 2-(Ethylthio)ethanol, Glycolamide, 2-Hydroxy-2-

- (hydroxymethyl)propanol, trans-2-Hydroxycyclohexanol, 2-Hydroxy-4-methylphenol, 2-(Hydroxymethyl)pyridine, 1-Hydroxymethyl-1-cyclohexanol, 2-Hydroxyhexanol, 2-Hydroxy-1-methoxypropane, 2-(Hydroxymethyl)imidazole, 3-Hydroxymethylpyrazole, trans-4-
- 5 Hydroxycyclohexanol, N-Acetyl-4-hydroxybutylamine, Hydroxycyclopentane, 2-(Methylsulfonyl)ethanol, 2-(Methylsulfinyl)ethanol, 4-(2-Hydroxyethyl)phenol, 2-(2-Hydroxyethyl)phenol, 2-Hydroxy-3-methylbutanol, 3-(N-Acetylamino)propanol, 3-(Diethylamino)propanol, 3-
- 10 (Dimethylamino)propanol, Allyl alcohol, 2-(Dimethylamino)ethanol, Glycerol, 2-Methoxyethanol, 2-(N-Acetylamino)ethanol, D-(Hydroxymethyl)pyrrolidine, 3-Hydroxypyrrolidine, 2-(Hydroxyethyl)benzene, 2-Hydroxyethyl-1-methylpyrrolidine, 2-Hydroxy-2-methyl-propanol, Cyclopropanol, Cyclohexanol, 3-
- Hydroxypropanol, 3-Ethoxypropanol, Propargyl alcohol, Ethyl glycolate, 2-Fluoroethanol, 3-(Dodecyloxy)propanol, 4-Hydroxybutanol, 5-Hydroxypentanol, 2-(Dimethylamino)ethanol, 2-(2-Hydroxyethoxy)ethanol, 1-(2-Hydroxyethyl)imidazolone, 2-(2-Hydroxyethoxy)ethylamine, Isopropanol, 2,2,2-Trifluoroethanol, 4-
- Nitrobenzyl alcohol, 3-Nitrobenzyl alcohol, 2-Methoxyethanol, 4(Hydroxyethyl)phenol, 4-(3-Hydroxypropyl)-1-sulfonamidobenzene,
 D,L-2-(Hydroxymethyl)tetrahydrofuran, Methyl lactate, 5Hydroxyhexanoic acid, methyl ester, 3-Methoxypropanol, 3Hydroxypiperidine, Pentanol, 4-Hydroxyheptane, 4-(2-Hydroxyethyl)-
- 1,2-dimethoxybenzene, 4-Hydroxymethyl-1,2-methylenedioxybenzene, 4-(Trifluoromethyl)benzyl alcohol, 4-(Methylthio)pheno, 2-(Hydroxymethyl)furan, 5-Hydroxypentan-2-one, 2-Hydroxy-3-methylbutanoic acid, methyl ester, 2-Hydroxy-3-phenyl-propanoic acid, ethyl ester, 1-(Hydroxymethyl)napthalene, 3-Phenylpropanol, 3,3-
- Dimethylbutanol, 3-(2-Hydroxyethyl)fluorobenzene, 4-Hydroxy-1-carboethoxypiperidine, (R)-2-(Hydroxymethyl)tetrahydrofuran, (S)-2-(Hydroxymethyl)tetrahydrofuran, (S)-2-Hydroxy-3-methylbutanol, (R)-2-Hydroxy-3-methylbutanol, (S)-2-Hydroxy-propanol, 3,4-Dihydroxytetrahydrofuran, 1,1,1,3,3,3-hexfluoroisopropanol, 2-

Fluorobenzyl alcohol, tert-Butanol, 2-Hydroxy-1-phenylethanol, iso-Butanol, 4-(2-Hydroxyethyl)fluorobenzene, 3-(Hydroxymethyl)toluene, 2-Chlorobenzyl alcohol, 2,4-Dichorobenzyl alcohol, sec-Butanol, R-2-Hydroxypropanol, Butanol, 4-Chlorobenzyl alcohol, 2-Ethoxyethanol, 2-(2-Hydroxyethyl)chlorobenzene, 2-(N-Methyl-N-phenylamino)ethanol, 3-(Trifluoromethyl)benzyl alcohol, 2-(Trifluoromethyl)benzyl alcohol, 2-(Hydroxyethyl)tetrahydrofuran, 4-Phenylbutanol, Nonyl alcohol, 2,6-Difluorobenzyl alcohol, 2-(Hydroxymethyl)thiophene, 2-(Hydroxyethyl)-1-methylpyrrole, 2-Hydroxy-3-methylbutane, 4-Hydroxymethyl-1,2-dichlorobenzene, 3-(Methylamino)propanol, 1,4-Difluorobenzyl alcohol, (2-Hydroxymethyl)furan,

EXAMPLE 29

N-Methyl nodulisporamide and 26-epi-N-methyl nodulisporamide

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To 1 mg nodulisporic acid in 1 mL dimethylformamide at room temperature was added 2 mg HCl•H2NMe, 2 mg N-hydroxybenzotriazole and 10 µL diisopropylethylamine to which was added 2 mg EDC•HCl. After 30 minutes, the reaction was quenched by addition of methanol and 1 drop glacial acetic acid. The solution was diluted with brine, extracted with ethyl acetate, dried with Na2SO4, filtered and concentrated under reduced pressure. The reaction was partially purified by preparative TLC (1 x 0.5 mm silica gel plate) using 6:3:1 EtOAc/acetone/methanol. N-Methyl nodulisporamide and 26-epi-N-methyl nodulisporamide were purified to homogeniety by reversed-phase HPLC using a 60 minute linear gradient from 25:75 to 100:0 acetonitrile/water. The purified products were characterized by ¹H NMR and mass spectrometry.

EXAMPLE 30 N-(n-Propyl)-nodulisporamide

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To 0.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 2 drops diisopropylethylamine, 5 mg H₂NCH₂CH₃, 3 mg N-hydroxylbenzotriazole and 3 mg PyBOP.

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After 30 minutes at room temperature, the reaction was quenched with 2 mL saturated NaHCO₃, extracted with ethyl acetate, dried (Na₂SO₄), filtered and concentrated in vacuo. The crude was partially purified by silica gel flash chromatography using 0.5:5:95 NH₄OH/MeOH/CHCl₃ as eluant followed by reversed-phase HPLC purification using 20:80 water/methanol as eluant. The product was characterized by ¹H NMR.

EXAMPLE 314-Morpholinyl-nodulisporamide

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To 1.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 1 drop morpholine and 2 mg N-hydroxybenzotriazole. 2 mg pyBOP was then added. After 1 hour at room temperature, the solution was filtered through 2 inches silica gel in a pipet without workup using ethyl acetate as eluant. The resultant solution was concentrated under reduced pressure and pure product was obtained following reversed-phase HPLC using 20:80 water/MeOH as eluant. The product was characterized by 1H NMR.

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EXAMPLE 32 N-(2-Hydroxyethyl)-nodulisporamide

To 0.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 2 drops diisopropylethylamine, 5 mg H₂NCH₂CH₂OH, 3 mg N-hydroxybenzotriazole and 3 mg PyBOP. After 30 minutes, the reaction was quenched with 2 mL saturated NaHCO₃, extracted with ethyl acetate, dried (Na₂SO₄), filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using 20:80 water/methanol as eluant and the product was characterized by ¹H NMR and mass spectrometry.

EXAMPLE 33

N-(1-Methoxycarbonyl-2-hydroxyethyl)-nodulisporamide

To 1.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 2 drops diisopropylethylamine, 5 mg HCl•H₂NCH(CH₂OH)CO₂Me, 3 mg N-hydroxybenzotriazole and 3 mg PyBOP. After 30 minutes, the reaction was quenched with 2 mL saturated NaHCO₃, extracted with ethyl acetate, dried (Na₂SO₄), filtered and concentrated in vacuo. Pure product was obtained following reversed-phase HPLC using 20:80 water/methanol as eluant and the product was characterized by ¹H NMR.

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EXAMPLE 34

Nodulisporamide and 31-amino-31,32-dihydro-nodulisporamide

To 1.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 1 drop NH4OH and 2 mg N-hydroxybenzotriazole. To this was added 3 mg PyBOP and the solution was stirred for 15 min. The reaction was quenched with 2 mL saturated NaHCO3, extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. Pure nodulisporamide was obtained following preparative TLC (1 x 0.5 mm silica gel) using 1:9 methanol/chloroform as eluant. Nodulisporamide was characterized by ¹H NMR and mass spectrometry. Also obtained from this reaction was 31-amino-31,32-dihydro-nodulisporamide.

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EXAMPLE 35

N-(Methoxycarbonylmethyl)-nodulisporamide

To 1.5 mg nodulisporic acid in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 2 mg N-hydroxybenzotriazole and 2 mg HCl•H2NCH2CO2Me. To this solution was added 2 mg PyBOP. After 30 min, the reaction was quenched with 2 mL saturated NaHCO3, extracted with ethyl acetate, dried using Na2SO4, filtered and concentrated under reduced pressure. Pure product was obtained following reversed-phase HPLC purification using

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17.5:82.5 water/methanol as eluant. The product was characterized by ¹H NMR and mass spectrometry.

EXAMPLE 36 N,N-Tetramethylene-nodulisporamide

To 125 mg nodulisporic acid in 10 mL methylene chloride at 0°C was added 0.18 mL diisopropylethylamine, 0.15 mL pyrrolidine followed by 108 mg PyBOP. After 5 minutes, the solution was warmed to room temperature. After 1.5 hours, the solution was poured in 25 mL saturated NaHCO3, extracted with methylene chloride, dried with Na2SO4, filtered and concentrated under reduced pressure. Pure N,N-tetramethylene-nodulisporamide was obtained following reversed-phase HPLC purification using 50:50 acetonitrile/water as eluant (isocratic for ten min), followed by a linear 30 minute gradient to 75:25 acetonitrile/water. Pure product (26 mg) was characterized by ¹H NMR and MS.

EXAMPLE 37

N-Ethyl 29,30-dihydro-20,30-oxa-nodulisporamide

To 1 mg Compound B in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 1 drop CH₃CH₂NH₂, 3 mg N-hydroxybenzotriazole and 3 mg PyBOP. After 15 minutes, the reaction was quenched with 2 mL saturated NaHCO₃, extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using 15:85 water/methanol as eluant and the purified product was characterized by ¹H NMR.

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EXAMPLE 38

N-(2-Hydroxyethyl)-29,30-dihydro-20,30-oxa-nodulisporamide

To 0.7 mg Compound B in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 1 drop HOCH₂CH₂NH₂, 3 mg N-hydroxybenzotriazole and 3 mg PyBOP.

After 15 minutes, the reaction was quenched with 2 mL saturated NaHCO3, extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using first 20:80 water/methanol then 15:85 water/methanol as eluant and the purified product was characterized by ¹H NMR.

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EXAMPLE 39

N-(2-Hydroxyethyl)-31-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-nodulisporamide

To 1 mg Compound C in 1 mL methylene chloride at room temperature was added 1 drop diisopropylethylamine, 1 drop HOCH2CH2NH2, 3 mg N-hydroxybenzotriazole and 3 mg PyBOP. After 15 minutes, the reaction was quenched with 2 mL saturated NaHCO3, extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated in vacuo. The crude was purified by reversed-phase HPLC using first 20:80 water/methanol as eluant and the purified product was characterized by ¹H NMR.

Example 40 N-tert-Butyl Nodulisporamide

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To a solution of 30 mg of nodulisporic acid in 3 mL methylene chloride at 0 °C was added 0.03 mL triethylamine and 12 mg N-hydroxybenzotriazole followed by 28 mg BOP reagent. The solution was stirred for 10 minutes and then 0.05 mL tert-butylamine was added. The solution was stirred overnight at 4 °C and then poured into 1/1 saturated sodium bicarbonate/brine, extracted with methylene chloride and the combined organic layers dried over sodium sulfate. The solids were removed by filtration and the solution concentrated to dryness under reduced pressure. The residue was partially purified by preparative TLC

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on silica gel (one 1000 micron plate) using 1/2 acetone/hexanes as eluant. Additional purification using HPLC (6/4 acetonitrile/water for 15 minutes, then a 45 minute linear gradient to 7/3 acetontrile/water) yielded pure product (17 mg). The purified product was characterized by proton NMR and MS (m/z: 735.7 (M+1)).

The general procedure of Example 40 was repeated using the appropriate amines listed in Table 3 below to provide the corresponding monosubstituted nodulisporamide compounds. These compounds were characterized by proton NMR and/or mass spectrometry (unless otherwise specified, m/z is for M+1).

Table 3: Monosubstituted Aliphatic Nodulisporamide Derivatives

15

Ex.	m/z	Amines	Rx
41	796.5	Aminoacetaldehyde diethyl acetal	
42	767.6	(2-Hydroxyethoxy)- ethylamine	CH2CH2OCH2CH2OH
43	792.5	4-(2-Aminoethyl)- morpholine	-CH ₂ CH ₂ -NO
44	790.4	1-(2-Aminoethyl)- piperidine	-CH ₂ CH ₂ -N
45	807.5	6-Amino-2-methylheptan- 2-ol	CH(CH3)(CH2)3C(CH3)2OH
46	737.5	3-Aminopropanol	(CH ₂) ₃ OH

	1		
47	751.5	4-Aminobutanol	(CH ₂) ₄ OH
48	765.6	5-Aminopentanol	(CH ₂)5OH
49	791.5	1-(2-Aminoethyl)- piperazine	-CH ₂ CH ₂ -N N
50	804.6	1-(3-Aminopropyl)-2- pyrrolidinone	-(CH ₂) ₃ -N
51	776.4	1-(2-Aminoethyl)- pyrrolidine	-(CH ₂) ₂ -N
52	751.4	2-Aminobutanol	CH(CH ₂ OH)CH ₂ CH ₃
53	750.5	tert-Butylhydrazine	NHC(CH3)3
54	718.3	Aminoacetonitrile	CH ₂ CN
55	779.6	6-Aminohexanol	(CH ₂) ₆ OH
56	806.8	4-(3-Aminopropyl)- morpholine	-CH ₂ CH ₂ CH ₂ -NO
57	737.4	3-Aminopropan-2-ol	CH ₂ CH(OH)CH ₃
58	765.4	2-Aminopentanol	CH(CH ₂ OH)CH ₂ CH ₂ CH ₃
59	777.7	1-Amino-1-cyclopentane- methanol	HOCH ₂
60		2-(Methylthio)ethylamine	CH2CH2SCH3
61	765.4	2-(Ethylthio)ethylamine	CH2CH2SCH2CH3
62	736.5	Glycineamide	CH ₂ CONH ₂
63	748.4	1-Aminopyrrolidine	-N
64		2-Amino-2- (hydroxymethyl)propanol	CH(CH ₃)(CH ₂ OH) ₂
65	777.6	trans-2- Aminocyclohexanol	НО
66	777.6	1-Amino-4-methyl- piperazine	−N N−CH ₃

67	766.5	2-(2-Aminoethylamino)-	CH ₂ CH ₂ NHCH ₂ CH ₂ OH
		ethanol	
68	791.4	1-Aminomethyl-	HO
		cyclohexan-1-ol	-CH ₂
69	779.4	2-Aminohexanol	CH(CH ₂ OH)(CH ₂) ₃ CH ₃
70	751.5	2-Amino-1-	CH(CH ₂ OCH ₃)CH ₃
		methoxypropane	
71	764.4	4-Aminomorpholine	_N_O
72	777.6	trans-4-Aminocyclohexan-	-{>-он
		1-ol	
73	739.4	2-Aminoethanethiol	(CH ₂) ₂ SH
74	750.5	4-Aminobutylamine	(CH ₂) ₄ NH ₂
75	764.4	2-Amino-4,5-	N—\
		dihydrothiazole	s
76	747.5	Aminocyclopentane	\triangle
77		2-(Methylsulfonyl)-	CH2CH2SO2CH3
		ethylamine	
78		2-(Methylsulfinyl)-	CH ₂ CH ₂ S(O)CH ₃
		ethylamine	
79	765.4	2-Amino-3-methylbutanol	CH(CH(CH ₃) ₂)CH ₂ OH
80	736.5	3-Aminopropylamine	(CH ₂) ₃ NH ₂
81	792.5	3-(Diethylamino)-	(CH ₂) ₃ N(CH ₂ CH ₃) ₂
		propylamine	
82	764.5	3-(Dimethylamino)-	(CH ₂) ₃ N(CH ₃) ₂
		propylamine	
83	723.5	O-Ethylhydroxylamine	OCH ₂ CH ₃
84	753.5	3-Amino-2-	CH ₂ CH(OH)CH ₂ OH
		hydroxypropanol	
85	709.4	O-Methylhydroxylamine	OCH3
86	737.4	2-Methoxyethylamine	CH2CH2OCH3

	564.4		
87			CH ₂ CH ₂ NHC(O)CH ₃
88	790.6	2-Aminoethyl-1-	H ₃ C-N
		methylpyrrolidine	
-			−CH ₂ CH ₂
89	751.5	2-Amino-2-methyl-	C(CH ₃) ₂ CH ₂ OH
<u></u>		propanol	
90	719.4	Cyclopropylamine	c-C3H5
91	760.5	Cyclohexylamine	c-C6H ₁₁
92	765.5	3-Ethoxypropylamine	(CH ₂) ₃ OCH ₂ CH ₃
93	719.5	Allylamine	CH2CH=CH2
94	789.5	2-Amino-2-	C(CH ₂ CH ₃)(CH ₂ OH) ₂
		(hydroxymethyl)butanol	2 - 3,2
95	717.5	Propargylamine	CH ₂ C≡CH
96	765.5	Glycine ethyl ester	CH2CO2CH2CH3
97	725.7	2-Fluoroethylamine	CH2CH2F
98	905.5	3-(Dodecyloxy)-	(CH ₂) ₃ O(CH ₂) ₁₁ CH ₃
	<u> </u>	propylamine	2,11012
99	751.0	2-(Dimethylamino)-	CH2CH2N(CH3)2
		ethylamine	
100	791.4	1-(2-Aminoethyl)-	Н
ļ		imidazolone	0=\(^N\)
			N/
101			−CH ₂ CH ₂
101	766.4	2-(2-Aminoethoxy)-	CH ₂ CH ₂ OCH ₂ CH ₂ NH ₂
100		ethylamine	
102		2,2,2-Trifluoroethylamine	CH ₂ CF ₃
103	780.5	Ethyl hydrazinoacetate	NHCH2CO2CH2CH3
104	763.5	D,L-2-(Aminomethyl)-	
		tetrahydrofuran	-CH ₂ -CO
105		1-Aminopiperidine	
			-N
106	765.6	D-Alanine methyl ester	CH(CH ₃)CO ₂ CH ₃

1.00	T=== =		
107	777.5	4-Amino-4-methyl-pentan-	C(CH ₃) ₂ CH ₂ C(O)CH ₃
		2-one	
108	837.6	Diethyl 2-aminomalonate	CH(CO ₂ CH ₂ CH ₃) ₂
109		5-Aminouracil	NH ONH
110	707.6	Ethylamine	CH ₂ CH ₃
111	807.8	Norleucine methyl ester	CH(CH ₂ CH ₂ CH ₃)CO ₂ CH ₃
112	751.7	3-Methoxypropylamine	CH2CH2CH2OCH3
113	745.5	1,1-Dimethylpropargylamine	C(CH ₃) ₂ C≡CH
114	749.7	Pentylamine	(CH2)4CH3
115	777.9	4-Aminoheptane	CH(CH ₂ CH ₂ CH ₃) ₂
116	763.8	Hexylamine	(CH ₂) ₅ CH ₃
117	776.8	cis-1,2- Diaminocyclohexane	H ₂ N
118	788.9	3-Aminoquinuclidine	
119	751.7	beta-Alanine	CH2CH2CO2H
120	793.5	L-Valine methyl ester	CH(CH(CH ₃) ₂)CO ₂ CH ₃
121		1-Amino-4-(2- Hydroxyethyl)piperazine	-N N·CH₂CH₂·OH
122	753.4	Aminooxyacetic acid	OCH2CO2H
123	834.5	4-Amino-1- carboethoxypiperidine	N-CO ₂ CH ₂ CH ₃
124	763.5	(R)-2-(Aminomethyl)- tetrahydrofuran	CH ₂ O
125	763.6	(S)-2-(Aminomethyl)- tetrahydrofuran	CH ₂ O

100	1-4-4		
126	1.000	L-Valinol	CH(CH(CH ₃) ₂)CH ₂ OH
127	765.7	D-Valinol	CH(CH(CH ₃) ₂)CH ₂ OH
128	737.7	L-Alaninol	CH(CH ₃)CH ₂ OH
129	737.6	D-Alaninol	CH(CH ₃)CH ₂ OH
130	721.7	Isopropylamine	CH(CH ₃) ₂
131	735.7	tert-butylamine	C(CH3)3
132	735.7	iso-Butylamine	(CH ₂)CH(CH ₃) ₂
133	735.5	sec-Butylamine	CH(CH ₃)CH ₂ CH ₃
134	737.6	(R)-3-Aminopropan-2-ol	CH ₂ CH(CH ₃)OH
135	735.6	n-Butylamine	(CH ₂) ₃ CH ₃
136	751.7	2-Ethoxyethylamine	(CH ₂) ₂ OCH ₂ CH ₃
137	787.7	2-Aminoethylcyclohexene	-CH ₂ CH ₂
138	813.7	1-Aminoadamantane	1-adamantyl
139	805.7	n-Nonylamine	(CH ₂) ₈ CH ₃
140	749.8	2-Amino-3-methylbutane	CH(CH ₃)CH(CH ₃) ₂
141	750.6	3-(Methylamino)- propylamine	(CH ₂) ₃ NHCH ₃
142	778.7	2-(Diethylamino)- ethylamine	(CH ₂) ₂ N(CH ₂ CH ₃) ₂
143	776.7	1-Amino-homopiperidine	-N

The general procedure of Example 40 was repeated using the amines listed in Table 4 below to provide the corresponding nodulisporamide compounds. These compounds were characterized by proton NMR and/or mass spectrometry (unless otherwise specified, m/z is for M+1).

Table 4: Nodulisporamide Derivatives

5

	, 		
Ex.	m/z	Amine	NRXRY
144	791.5	1-(2-Aminoethyl)- piperazine	N-CH ₂ CH ₂ -NH ₂
145	776.6	4-Aminomethylpiperidine	-N-CH ₂ NH ₂
146	765.4	Thiomorpholine	$-$ N $ _{\mathbf{S}}$
147	759.4	Diallylamine	N(CH ₂ CH=CH ₂) ₂
148	737.4	2-(Methylamino)ethanol	N(CH ₃)CH ₂ CH ₂ OH
149	795.4	Diisopropanolamine	N(CH ₂ CH(CH ₃)OH) ₂
150	763.5	L-2-(Hydroxymethyl)- pyrrolidine	HO N
151	763.5	D-2-(Hydroxymethyl)- pyrrolidine	HO
152	749.5	3-Hydroxypyrrolidine	~ ОН
153	732.7	Methylaminoacetonitrile	N(CH3)CH2C≡N
154		4-(2-hydroxyethyl)- piperazine	-NN-CH₂CH₂OH
155	777.7	4-Ethylpiperazine	-NN·CH₂CH₃
156	721.5	N-Ethylmethylamine	N(CH ₃)CH ₂ CH ₃
157	735.6	N-(Methyl)isopropylamine	N(CH3)CH(CH3)2

			
158		N-Methylpropylamine	N(CH ₃)CH ₂ CH ₂ CH ₃
159	749.5	N-Methylbutylamine	N(CH ₃)CH ₂ CH ₂ CH ₂ CH ₃
160	765.7	N-Ethyl-2-methoxyethyl-	N(CH2CH3)CH2CH2OCH3
		amine	
161	751.7	N-Methyl-2-methoxyethyl-	N(CH ₃)CH ₂ CH ₂ OCH ₃
		amine	
162	749.7	N-Ethylpropylamine	N(CH ₂ CH ₃)CH ₂ CH ₂ CH ₃
163	751.5	Tetrahydrothiazole	N
) S
164	767.8	Diethanolamine	N(CH ₂ CH ₂ OH) ₂
165	763.8	3-Hydroxypiperidine	(617261172
105	705.0	3-Hydroxypiperidille	-N
-			ОН
166	763.9	4-Hydroxypiperidine	-N—OH
			-N OH
167	749.6	N-(Ethyl)isopropylamine	N(CH2CH3)CH(CH3)2
168	747.8	Piperidine	
			-N
169	735.8	Diethylamine	N(CH ₂ CH ₃) ₂
170	762.7	4-Methylpiperazine	
ĺ			−N N−CH ₃
171	767.6	Tetrahydrothiazole-S-	
- / -	707.0	oxide	N S+0
1.50	-		
172	791.7	Dibutylamine	N(CH ₂ CH ₂ CH ₃) ₂
173	745.7	1,2,3,6-Tetrahydropyridine	-N
			'`
174	790.8	3-(Carboxamido)piperidine	
			-N >
		İ	CONH ₂
			0011112

			
175	819.6	3-(Carboethoxy)piperidine	-N CO ₂ CH ₂ CH ₃
176	761.6	Hexamethyleneimine	
	701.0	Trexamenty tenenimic	\N\
177	820.7	1-(Carboethoxy)piperazine	−NN-CO₂CH₂CH₃
178	819.7	Dipentylamine	N(CH2CH2CH2CH3)2
179	775.6	Heptamethyleneimine	- N
180	787.6	Octahydroindole	
181	760.5	4,5-Dihydro-5,5-	N-10
		dimethylimidazole	CH ₃ CH ₃
182	707.5	Dimethylamine	N(CH3)2
183	763.7	Dipropylamine	N(CH2CH2CH3)2
184	761.7	2-Methylpiperidine	-N H ₃ C
185	779.5	2-(Butylamino)ethanol	N((CH ₂) ₂ CH ₃)CH ₂ CH ₂ OH
186	731.7	Methylpropargylamine	N(CH ₃)CH ₂ C≡CH
187	854.7	1-(4-Methoxyphenyl)- piperazine	-NN-OCH3
188	931.9	Dinonylamine	N((CH ₂)8CH ₃) ₂
189	903.8	Dioctylamine	N((CH ₂)7CH ₃) ₂

100	10455		
190	815.7	, , , , , , , , , , , , , , , , , , ,	CH ₃
		aza[3.2.1]bicyclooctane	
		1	CH ₃
<u></u>			CH ₃
191	750.7	N,N'-Dimethylethylene-	N(CH ₃)(CH ₂) ₂ NHCH ₃
		diamine	
192	750.6	3-(Methylamino)-	N(CH ₃)(CH ₂) ₃ NH ₂
<u></u>		propylamine	_
193	813.7	L-2-Amino-3-	NHCH(CH2OH)CH2Ph
L		phenylpropanol	
194	785.6	2-Amino-4-methylphenol	NHPh(2-OH,4-CH3)
195		4-Aminobenzylamine	NHCH ₂ Ph(4-NH ₂)
196	789.4	4-Chloroaniline	NHPh(4-Cl)
197	799.5	4-(2-Hydroxyethyl)aniline	NHPh(4-CH2CH2OH)
198	799.5	2-(2-Hydroxyethyl)aniline	NHPh(2-CH ₂ CH ₂ OH)
199	783.4	2-Phenylethylamine	NHCH2CH2Ph
200	785.4	2-(Hydroxymethyl)aniline	NHPh(2-CH ₂ OH)
201	798.8	3-(Dimethylamino)aniline	NHPh(3-N(CH3)2
202	835.1	4-(Sulfonylamido)aniline	NHPh(4-SO ₂ NH ₂)
203		Phenylhydrazine	NHNHPh
204	798.4	2-Carboxamidoaniline	NHPh(2-CONH2)
205	799.8	4-(Aminoethyl)phenol	NHCH2CH2Ph(4-OH)
206	884.5	4-(3-Aminopropyl)-1-	NHCH2CH2Ph(4-SO2NH2)
		sulfonamidobenzene	2 == 2= 11(+ 3 32: 1112)
207	770.5	2-Aminoaniline	NHPh(2-NH2)
208	883.7	L-Leucine benzyl ester	NHCH(CH2CH(CH3)2)CO2
			CH ₂ Ph
209	888.5	4-(tert-butyl)benzyl-	NHSO ₂ CH ₂ Ph(4-C(CH ₃) ₃)
		sulfonamide	2 - 2 ((
210	833.6	Benzylsulfonamide	NHSO2CH2Ph
211	788.7	2-Fluorophenylhydrazine	NHNHPh(2-F)

212 843.8 4-(2-Aminoethyl)-1,2- dimethoxybenzene NHCH ₂ CH ₂	OMe
NHOH ₂ OH ₂	\
212 967 5 I Proling have 1	≻ OMe
212 967 5 I Decline bereat	
213 867.5 L-Proline benzyl ester	
PhCH ₂ O ₂ C	
214 813.8 4-Aminomethyl-1,2-	
methylenedioxybenzene HN.	
~ 0	
215 837.5 4-(Trifluoromethyl)- NHCH ₂ Ph(4-CF ₃)	
benzylamine	
216 882.6 1-((3,4-methylenedioxy)-	$\mathbf{P}^{\mathbf{Q}}$
benzyl)piperazine	
217 862.7 3-(Benzyloxy)aniline NHPh(4-OCH ₂ Ph)	
218 801.4 4-(Methylthio)aniline NHPh(4-SCH3)	
219 855.5 L-Phenylalanine ethyl ester NHCH(CH2Ph)CO2C	H ₂ CH ₃
220 841.4 D-Phenylalanine methyl NHCH(CH2Ph)CO2C	H 3
ester	
221 799.4 4-(Methoxy)benzylamine NHCH2Ph(4-OCH3)	
222 819.5 1- NHCH2-1-naphthyl	
(Aminomethyl)napthalene	
223 792.4 1,2,3,4-Tetrahydro-	
isoquinoline	
224 821.8 3-Amino-2-hydroxy- HO	
- Julius 2 my drony	
napthalene H	
225 801.7 3-(2-Aminoethyl)- NHCH2CH2(3-F)Ph	
fluorobenzene	
226 823.7 4-Phenylpiperazine N-Ph	
IN I	
227 814.7 D-Phenylalaninol NHCH(CH ₂ Ph)CH ₂ O	Н

000	1000 6		
228	8 838.6	1-(o-Tolyl)piperazine	-N
<u> </u>			H ₃ C
229	847.6	Spiro(1H-indene-1,4'-piperidine)	
230	773.6	4-Fluoroaniline	NHPh(4-F)
231	787.5	2-Fluorobenzylamine	NHCH ₂ Ph(2-F)
232	799.7	2-Amino-1-phenylethanol	NHCH2CH(Ph)OH
234	801.8	4-(2-Aminoethyl)-1-	NHCH2CH2Ph(4-F)
		fluorobenzene	,
235	829.5	4-(2-Amino-2-	NHC(CH ₃) ₂ CH ₂ Ph(3-F)
		methylpropyl)-1-	
		fluorobenzene	
236	791.7	3,4-Difluoroaniline	NHPh(3,4-diF)
237	783.7	3-(Aminomethyl)toluene	NHCH ₂ Ph(3-CH ₃)
238	784.5	3-Methylphenylhydrazine	NHNH(3-CH ₃)Ph
239	803.5	2-Chlorobenzylamine	NHCH ₂ Ph(2-Cl)
240	838.8	2,4-Dichorobenzylamine	NHCH2Ph(2,4-diCl)
241	782.7	4-Methylphenylhydrazine	NHNHPh(4-CH3)
242	803.8	4-Chlorobenzylamine	NHCH2Ph(4-Cl)
243	797.7	3-Phenylpropylamine	NH(CH2)3Ph
244	817.6	4-(2-Aminoethyl)-1-	NHCH2CH2Ph(4-Cl)
		chlorobenzene	_ ,
245	893.8	1-(m-Trifluoromethyl	
		phenyl)piperazine	-N N
			CF₃
246	852.6	1-(2,3-Dimethylphenyl)	
		piperazine	-N N
			H ₃ C CH ₃
		·	1130 0113

	-1-		
247	812.7	N-Methyl-N-phenyl-	NHCH2CH2N(CH3)Ph
		ethylenediamine	
248	837.6	3-(Trifluoromethyl)-	NHCH ₂ Ph(3-CF ₃)
		benzylamine	
249	837.7	2-(Trifluoromethyl)-	NHCH ₂ Ph(2-CF ₃)
		benzylamine	
250		1-(4-Methoxyphenyl)-	
		piperazine	N N OCH3
251	795.7	2-Aminoindane	
			HN
252	843.6	9-Aminofluorene	HN
253	811.7	4-Phenylbutylamine	NH(CH ₂) ₄ Ph
254	827.8	(R,R)-2-Methylamino-3-	N(CH ₃)CH(CH ₃)CH(CH ₃)Ph
		phenylbutane	
255	827.8	(S,S)-2-Methylamino-3-	N(CH ₃)CH(CH ₃)CH(CH ₃)Ph
		phenylbutane	
256	825.9	Benzylbutylamine	N(CH ₂ Ph)(CH ₂) ₃ CH ₃
257	785.6	O-Benzylhydroxylamine	NHOCH ₂ Ph
258	805.5	2,6-Difluorobenzylamine	NCH ₂ Ph(2,6-diF)
259	920.9	1-(2-(o-Trifluoromethyl-	F ₃ C
		phenyl)ethyl)piperazine	$\left - \stackrel{\wedge}{N} \right $ $\left - N \right $ $\left - CH_2 CH_2 \right $
			1 0.20.2
260	797.7	(S)-N,alpha-	N(CH3)CH(CH3)Ph
		Dimethylbenzylamine	
261	783.7	(S)-alpha-	NHCH(CH3)Ph
		Methylbenzylamine	
262	797.6	Methyl benzyl amine	N(CH ₃)CH ₂ Ph
263		4-Aminomethyl-1,2-	NHCH ₂ Ph(3,4-diCl)
		dichlorobenzene	

	1 = 22 =		
264	1 783.7	1 (- >F	N(CH ₃)CH(CH ₃)Ph
		Methylbenzylamine	
265	873.8	1-Benzylamino-2-	N(CH ₂ Ph)CH ₂ CH ₂ Ph
		phenylethane	
266	784.6	Benzylhydrazine	NHNHCH2Ph
267	805.7	2,4-Difluorobenzylamine	NHCH ₂ Ph(2,4-diF)
268	838.8	2,5-Dichlorophenyl-	NHNHPh(2,5-diCl)
		hydrazine	(2,5 (2.61)
269	787.7	3-Fluorobenzylamine	NHCH ₂ Ph(3-F)
270	795.5	1-Aminoindane	HN
271	859.8	1,2-Diphenylethylamine	NHCH(Ph)CH2Ph
272	801.8	3,4-Dihydroxybenzylamine	
273	829.7	2,4-Dimethoxy-	NHCH2Ph(3,4-diOCH3)
		benzylamine	2 = (0, 1 = 0 = 0 = 0, 1
274	783.8	N-Benzylmethylamine	N(CH3)CH2Ph
275	797.7	N-Benzylethylamine	N(CH2CH3)CH2Ph
276		(R)-N,alpha-	N(CH3)CH(CH3)Ph
		Dimethylbenzylamine	3,(3,
277	770.5	3-(Aminomethyl)pyridine	HN, CEN
		7 /7 / / / / / / / / / / / / / / / / /	CH ₂ -(
-			
278	745.9	3-Amino-1,2,4-triazole	H N
			HN N
279	757.4	2-Aminopyrimidine	HN
)=N
			N
280	784.6	2-(2-Aminoethyl)pyridine	HN
		3 /13	>=N

	,		
281	787.5	1-(3-Aminopropyl)- imidazole	HN — (CḤ₂) ₃ N
		imidazoie	
282	770.6	4-(Aminomethyl)pyridine	-NHCH ₂
283	757.4	2-Aminopyrazine	HN—N=
284		3-Amino-1,2,4-triazine	HN-⟨N-N
285		5-Amino-3- hydroxypyrazole	HO NH
286		2-Amino-3- hydroxypyridine	HO N=
287		4-Amino-5- carboxamidoimidazole	H N CONH ₂
288	770.4	2-(Aminomethyl)pyridine	-NHCH ₂ —
289	751.5 M+Li	2-Aminoimidazole	HN HN
290	745.4	3-Aminopyrazole	HN N-N
291	795.2	6-Aminobenzopyrazole	HN N-N
292	797.5	4-Amino-1,2,4-triazole	HN N N

293	3	2-Amino-4,5- dihydrothiazole	HN S
294	762.4	2-Aminothiazole	HN S
295	795.4	5-Aminobenzopyrazole	HN
296	761.6	3,5-Diamino-1,2,4-triazole	HN N N N NH ₂
297	825.7	1-(2-Pyridyl)piperazine	$N \longrightarrow N \longrightarrow N$
298	798.7	4-(Ethylaminomethyl)- pyridine	N-CH ₂ -N CH ₂ CH ₃
299	1032. 7	L-Tryptophan-1,1- diphenylmethylamide	HN CONHCH(Ph) ₂
300		2-(Aminomethyl)thiophene	HN S
301		2-(2-Aminoethyl)-1- methylpyrrole	HN CH ₃
302	759.5	2-(Aminomethyl)furan	HNO

EXAMPLE 303

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General Procedure for the Preparation of Additional Amide Derivatives of Nodulisporic Acid

To a solution of 30 mg of nodulisporic acid in 3 mL methylene

chloride at 0 °C add 0.03 mL triethylamine and 12 mg Nhydroxybenzotriazole followed by 28 mg BOP reagent. Stir the solution
for 10 minutes and then add 50 mg of amine selected from Table 5. Stir
the solution overnight at 4 °C and then pour into 1/1 saturated sodium
bicarbonate/brine, extract with methylene chloride and dry the combined
organic layers over sodium sulfate. Remove the solids by filtration and
concentrate the solution to dryness under reduced pressure. Pure product
may be obtained by flash chromatography or preparative TLC on silica
gel or reversed-phase liquid chromatography. The purified product may
be characterized by proton NMR and mass spectrometry.

Table 5: Amines for the Preparation of Additional Nodulisporamide Derivatives

15

N-Methyl-2,2,2-trifluoroethylamine, 2,2,3,3,3-Pentafluoropropylamine,
N-Methyl-2,2,3,3,3-pentafluoropropylamine, 1,1,1,3,3,3Hexafluoroisopropylamine, 2-Difluoro-3-Methoxy-1-methylpropylamine, N-Methyl-1,1,1,3,3,3-hexafluoroisopropylamine, 1,1,1Trifluoromethylpropylamine, 2-(3,3,3-Trifluoromethyl)propylamine, NMethyl-1,1,1,3,3,3-hexafluoroisopropylamine, Di-(2,2,2-

- trifluoroethyl)amine, N-(2-Methoxyethyl)-2,2,2-trifluoroethylamine, 2-Methoxy-1-methyl-ethylamine, 3-Methoxy-1-methyl-propylamine, 2-Methoxy-1-methyl-ethylamine, N-Methyl-2-methoxy-1-benzyl-ethylamine, 1-Methoxymethyl-3-methyl-butylamine, Methylsulfonamide, Isopropylsulfonamide, Ethylsulfonamide, Benzylsulfonamide, sec-
- Butylsulfonamide, N-Methyl-ethylsulfonamide, N,1,1-Trimethyl-propargylamine, N-Ethyl-1,1-dimethyl-propargylamine, N,1-Dimethyl-propargylamine, 1-Methyl-propargylamine, 1-Trifluoromethylpropargylamine, N,1,1-Trimethyl-propargylamine, N-Ethyl-1,1-dimethyl-propargylamine, N,1-Dimethyl-propargylamine,

- N,1,1-Trimethyl-propargylamine, 1-Methyl-propargylamine, 1-Trifluoromethylpropargylamine, N-Ethylpropargylamine, N-(2-Methoxyethyl)propargylamine, 1-Amino-2-pentyne, 1-Amino-3-pentyne, 1-Amino-4-pentyne, 1-Methylamino-3-
- pentyne, 1-Methylamino-4-pentyne, 1-Ethylamino-4-pentyne, 1-Trifluoromethylamino-2-pentyne, 1-Trifluoromethylamino-3-pentyne, 1-Trifluoromethylamino-4-pentyne, N-(2-Methoxyethyl)-2-amino-1,1-dimethyl-2-butyne, 1-Amino-2-butyne, 1-Amino-3-butyne, N-Methylamino-2-butyne, 1-Ethylamino-3-
- butyne, 2-(Aminomethyl)dioxane, 2-(2-Aminoethyl)dioxane, 2-(3-Aminopropyl)dioxane, 2-(2-Aminopropyl)dioxane, 2-(Methylaminomethyl)dioxane, 2-(1-Aminoethyl)dioxane, 2-Aminomethyl-2H-tetrahydropyran, 2-(2-Aminopropyl)-2H-tetrahydropyran, 2-(2-Aminopropyl)-2H-tetrahydropyran, 2-(2-
- Aminopropyl)-2H-tetrahydropyran, 2-(2-Aminoethyl)-5-ethyl-2H-tetrahydropyran, 2-Methylaminomethyl-2H-tetrahydropyran, 2-(1-Aminoethyl)-2H-tetrahydropyran, 2-(2-Aminopropyl)tetrahydrofuran, 2-Aminomethyl-5-ethyl-tetrahydrofuran, 2-Methylaminomethyl-tetrahydrofuran, 2-(Ethylaminomethyl)tetrahydrofuran, 2-(1-
- Aminoethyl)tetrahydrofuran, 4-(Methoxymethyl)benzylamine, 4-(2-Methoxyethyl)benzylamine, 4-(Ethoxymethyl)benzylamine, 4-(Acetoxyoxymethyl)benzylamine, 3-(Dimethylaminomethyl)benzylamine, 4-(Sulfonamidomethyl)benzylamine, 2-Chloro-6-fluoro-benzylamine, 3-
- Chloro-4-fluoro-benzylamine, 2-Chloro-4-fluoro-benzylamine, 3,5-Difluoro-benzylamine, 2,4-Difluoro-benzylamine, Pentafluorobenzylamine, 4-Methoxy-2,3,5,6-tetrafluorobenzylamine, 4-(Methyl)benzylamine, Benzylamine, 4-(Ethyl)benzylamine, 4-(Ethoxy)benzylamine, 4-(Isopropyl)benzylamine, 4-
- (Isobutyl)benzylamine, 4-(Isopropropoxy)benzylamine, 4-(Isobutoxy)benzylamine, 4-(Allyl)benzylamine, 4-(Allyl)benzylamine, 4-(Allyl)benzylamine, 4-(Trifluoromethoxy)benzylamine, 4-(2,2,2-trifluoroethoxy)benzylamine, 3,4-Ethylenedioxybenzylamine, 4-Methoxymethyl-2-chloro-

phenethylamine, 4-(2-Methoxyethyl)phenethylamine, 4-(Ethoxymethyl)phenethylamine, 4-(Acetoxyoxymethyl)phenethylamine. 3-(Dimethylaminomethyl)phenethylamine, 1-Phenyl-2,2,2trifluoroethylamine, 4-(Trifluoromethoxy)aniline, 4-Methoxyaniline, 4-Ethoxyaniline, 3-Chloro-4-fluoro-aniline, 4-Chloro-2-fluoro-aniline, 4-5 (Acetoxy)aniline, 4-(Butoxy)aniline, 3-Chloroaniline, 4-(Methylthio)aniline, 5-(Aminomethyl)benzofuran, 5-(Methylaminomethyl)benzofuran, 4-(1-Aminoethyl)benzofuran, 5-(2-Aminoethyl)benzofuran, 5-Aminomethyl-2,3-dihydro-benzofuran, 5-Methylaminomethyl-2,3-dihydro-benzofuran, 4-1-Aminoethyl-2,3-10 dihydro-benzofuran, 5-2-Aminoethyl-2,3-dihydro-benzofuran, 5-Aminomethyl-2H-tetrahydrobenzopyran, 5-Methylaminomethyl-2Htetrahydrobenzopyran, 4-1-Aminoethyl-2H-tetrahydrobenzopyran, 5-2-Aminoethyl-2H-tetrahydrobenzopyran, 5-Aminomethyl-2Htetrahydrobenzopyran, 5-Methylaminomethyl-2H-tetrahydrobenzopyran, 15 4-(1-Aminoethyl)-2H-tetrahydrobenzopyran, 5-(2-Aminoethyl)-2Htetrahydrobenzopyran, 5-Aminomethyl-benzo-1,4-dioxane, 5-Methylaminomethyl-benzo-1,4-dioxane, 4-1-Aminoethyl-benzo-1,4dioxane, 5-2-Aminoethyl-benzo-1,4-dioxane, 5-Aminomethyl-benzo-1,4dioxane, 5-Methylaminomethyl-benzo-1,4-dioxane, 4-(1-Aminoethyl)-20 benzo-1,4-dioxane, 5-(2-Aminoethyl)-benzo-1,4-dioxane, 3-Amino-5methoxy-thiophene, 2-Amino-5-chloro-thiophene, 2-(2-Aminoethyl)thiophene, 2-(3-Aminopropyl)thiophene, 3-(3-Aminopropyl)thiophene, 3-(2-Methylaminoethyl)thiophene, 2-Chloro-3-(2-aminoethyl)-thiophene, 2-Aminoethyl-4-methoxy-thiophene, 2-25 Amino-3-ethyl-thiophene, 2-(Methylaminomethyl)thiophene, 3-(Aminomethyl)thiophene, 2-(2-Aminoethyl)-4-methoxy-thiophene, 1-(Aminomethyl)tetrazole, 1-(1-Aminoethyl)tetrazole, 1-(3-Aminopropyl)tetrazole, 5-Amino-3-methyl-isoxazole, 3-Aminopyridine, 4-Aminomethylthiazole, 2-(2-Aminoethyl)pyrazine, 2-(1-30 Aminoethyl)imidazole, 2-(Aminomethyl)isoxazole, 3-(2-

Aminoethyl)pyrazole, 2-(Aminomethyl)-1,3,4-thiadiazole.

General Procedure for Synthesis of Amide Derivatives of Compounds B and C

To a solution of 30 mg of compound B or C in 3 mL methylene chloride at 0 °C add 0.03 mL triethylamine and 12 mg N-hydroxybenzotriazole followed by 28 mg BOP reagent. Stir the solution for 10 minutes and then add 50 mg of an amine selected from Table 6. Stir overnight at 4 °C and then at room temperature for 2 hours. Pour the solution into 1/1 saturated sodium bicarbonate/brine. Extract the solution with methylene chloride and dry the combined organic layers over sodium sulfate. Remove the solids by filtration and concentrate the solution under reduced pressure. Pure product may be obtained following purification by flash chromatography, preparative TLC or reversed-phase liquid chromatography. Products may be characterized by proton NMR and/or mass spectrometry.

TABLE 6: Additional Amide Derivatives of Compounds B and C

- 2-(2-Hydroxyethoxy)ethylamine, 4-(2-Aminoethyl)morpholine, 1-(2-Aminoethyl)piperidine, 6-Amino-2-methylheptan-2-ol, 3-(Aminomethyl)pyridine, 3-Aminopropanol, 4-Aminobutanol, 5-Aminopentanol, 2-(2-Aminoethyl)piperidine, 1-(3-Aminopropyl)-2-pyrrolidinone, 1-(2-Aminoethyl)pyrrolidine, 2-Aminobutanol, 4-
- 25 (Aminomethyl)pyridine, 2-Aminopyrazine, tert-Butylhydrazine, 6-Aminohexanol, 4-(3-Aminopropyl)morpholine, 3-Aminopropan-2-ol, 2-Aminopentanol, 1-Amino-1-hydroxymethyl-cyclopentane, 2-(Methylthio)ethylamine, 2-(Ethylthio)ethylamine, Thiomorpholine, 4-Amino-5-carboxamidoimidazole, 1-Aminopyrrolidine, 2-Amino-2-
- hydroxymethyl-propanol, trans-2-Aminocyclohexan-1-ol, 4-Aminobenzylamine, 2-(Aminomethyl)pyridine, 1-Aminomethyl-cyclohexan-1-ol, 2-Amino-1-methoxypropane, 2-Aminoimidazole, 4-Aminomorpholine, trans-4-Aminocyclohexan-1-ol, 4-Amino-1,2,4-triazole, 2-Amino-4,5-dihydrothiazole, 2-(Methanesulfonyl)ethylamine,

- 2-(Methanesulfinyl)ethylamine, 4-(2-Hydroxyethyl)aniline, 2-(2-Hydroxyethyl)aniline, 2-Amino-3-methylbutanol, Diallylamine, 2-(Methylamino)ethanol, O-Ethylhydroxylamine, 3-Amino-2-hydroxypropanol, O-Methylhydroxylamine, L-
- 5 (Hydroxymethyl)pyrolidine, 2-Methoxyethylamine, N-Acetylethylenediamine, D-(Hydroxymethyl)pyrrolidine, 3-Hydroxypyrolidine, 2-(Aminoethyl)benzene, 2-Amino-2-methyl-propanol, Cyclohexylamine, 3-Ethoxypropylamine, Allylamine, 2-Amino-2-hydroxymethyl-butanol, Propargylamine, 2-Fluoroethylamine,
- 3-(Dimethylamino)aniline, 2-Dimethylaminoethanol, 4-(2-hydroxyethyl)piperazine, 4-Ethylpiperazine, N-Ethylmethylamine, N-(Methyl)isopropylamine, 2,2,2-Trifluoroethylamine, N-Methylpropylamine, N-Methylbutylamine, N-Ethyl-2-methoxyethylamine, 4-(Aminoethyl)phenol, N-Methyl-2-
- 15 methoxyethylamine, N-Ethylpropylamine, D,L-2(Aminomethyl)tetrahydrofuran, 1-Aminopiperidine, D-Alanine methyl
 ester, 3,5-Diamino-1,2,4-triazole, Benzylsulfonamide, 4-Amino-4methyl-pentan-2-one, 5-Aminouracil, Ethylamine, Norleucine methyl
 ester, 3-Methoxypropylamine, 3-Hydroxypiperidine, 4-
- Hydroxypiperidine, 1,1-Dimethylpropargylamine, N(Ethyl)isopropylamine, Pentylamine, Piperidine, 2Fluorophenylhydrazine, Hexylamine, Diethylamine, 4-(2-Aminoethyl)1,2-dimethoxybenzene, 1-(2-Pyridyl)piperazine, 4-Methylpiperazine, 4(2-Hydroxyethyl)morpholine, 4-Aminomethyl-1,2-
- 25 methylenedioxybenzene, 1-((3,4-methylenedioxy)benzyl)piperazine, 4(Ethylaminomethyl)pyridine, L-Valine methyl ester, D-Phenylalanine
 methyl ester, 4-(Methoxy)benzylamine, 1-Amino-4-(2hydroxyethyl)piperazine, 1,2,3,6-Tetrahydropyridine, 3-(2Aminoethyl)fluorobenzene, 1-Phenylpiperazine, 4-Amino-1-
- carboethoxypiperidine, 1-(Carboethoxy)piperazine, (R)-2(Aminomethyl)tetrahydrofuran, (S)-2-(Aminomethyl)tetrahydrofuran, LValinol, D-Valinol, L-Alaninol, D-Phenylalaninol, 3,4Dihydroxytetrahydrofuran, D-Alaninol, 2-Fluorobenzylamine, 4Fluoroaniline, Isopropylamine, tert-Butylamine, iso-Butylamine, 4-(2-

Aminoethyl)fluorobenzene, 4,5-Dihydro-5,5-dimethylimidazole, sec-Butylamine, Dimethylamine, (R)-3-Aminopropan-2-ol, Di-n-propylamine, n-Butylamine, 2-Methylpiperidine, 4-Chlorobenzylamine, 3-Phenylpropylamine, 2-Ethoxyethylamine, Methylpropargylamine, 2-

- 5 (Trifluoromethyl)benzylamine, 4-Phenylbutylamine, O-Benzylhydroxylamine, 2,6-Difluorobenzylamine, 2-(Aminomethyl)thiophene, 2-(2-Aminoethyl)-1-methylpyrrole, (S)-N,alpha-Dimethylbenzylamine, 2-Amino-3-methylbutane, (S)-alpha-Methylbenzylamine, 1-Methylamino-2-phenylethane, 3,4-
- Dichlorobenzylamine, 1,4-Difluorobenzylamine, 2-(Aminomethyl)furan, 3-Fluorobenzylamine, 2,4-Dimethoxybenzylamine, N-Benzylmethylamine, N-Ethylbenzylamine, N-Methyl-2,2,2-trifluoroethylamine, 2,2,3,3,3-Pentafluoropropylamine, N-Methyl-2,2,3,3,3-pentafluoropropylamine, 1,1,1,3,3,3-Hexafluoroisopropylamine
- 15 , 2-Difluoro-3-Methoxy-1-methyl-propylamine, N-Methyl-1,1,1,3,3,3-hexafluoroisopropylamine, 1,1,1-Trifluoromethylpropylamine, 2-(3,3,3-Trifluoromethyl)propylamine, N-Methyl-1,1,1,3,3,3-hexafluoroisopropylamine, Di-(2,2,2-trifluoroethyl)amine, N-(2-Methoxyethyl)-2,2,2-trifluoroethylamine, 2-Methoxy-1-methyl-
- ethylamine, 3-Methoxy-1-methyl-propylamine, 2-Methoxy-1-methylethylamine, N-Methyl-2-methoxy-1-benzyl-ethylamine, 1-Methoxymethyl-3-methyl-butylamine, Methylsulfonamide, Isopropylsulfonamide, Ethylsulfonamide, Benzylsulfonamide, sec-Butylsulfonamide, N-Methyl-ethylsulfonamide, N,1,1-Trimethyl-
- propargylamine, N-Ethyl-1,1-dimethyl-propargylamine, N,1-Dimethyl-propargylamine, 1-Methyl-propargylamine, 1-Trifluoromethylpropargylamine, N,1,1-Trimethyl-propargylamine, N-Ethyl-1,1-dimethyl-propargylamine, N,1-Dimethyl-propargylamine, N,1,1-Trimethyl-propargylamine, 1-Methyl-propargylamine, 1-
- Trifluoromethylpropargylamine, N-Ethylpropargylamine, N-(2-Methoxyethyl)propargylamine, 1-Amino-2-pentyne, 1-Amino-3-pentyne, 1-Amino-4-pentyne, 1-Methylamino-2-pentyne, 1-Methylamino-3-pentyne, 1-Methylamino-4-pentyne, 1-Ethylamino-4-pentyne, 1-Trifluoromethylamino-2-pentyne, 1-Trifluoromethylamino-3-pentyne, 1-

- Trifluoromethylamino-4-pentyne, N-(2-Methoxyethyl)-2-amino-1,1dimethyl-2-butyne, 1-Amino-2-butyne, 1-Amino-3-butyne, N-Methylamino-2-butyne, N-Methylamino-3-butyne, 1-Ethylamino-3butyne, 2-(Aminomethyl)dioxane, 2-(2-Aminoethyl)dioxane, 2-(3-
- Aminopropyl)dioxane, 2-(2-Aminopropyl)dioxane, 2-5 (Methylaminomethyl)dioxane, 2-(1-Aminoethyl)dioxane, 2-Aminomethyl-2H-tetrahydropyran, 2-(2-Aminoethyl)-2Htetrahydropyran, 2-(3-Aminopropyl)-2H-tetrahydropyran, 2-(2-Aminopropyl)-2H-tetrahydropyran, 2-(2-Aminoethyl)-5-ethyl-2H-
- tetrahydropyran, 2-Methylaminomethyl-2H-tetrahydropyran, 2-(1-10 Aminoethyl)-2H-tetrahydropyran, 2-(2-Aminopropyl)tetrahydrofuran, 2-Aminomethyl-5-ethyl-tetrahydrofuran, 2-Methylaminomethyltetrahydrofuran, 2-(Ethylaminomethyl)tetrahydrofuran, 2-(1-Aminoethyl)tetrahydrofuran, 4-(Methoxymethyl)benzylamine, 4-(2-
- Methoxyethyl)benzylamine, 4-(Ethoxymethyl)benzylamine, 4-15 (Acetoxyoxymethyl)benzylamine, 3-(Dimethylaminomethyl)benzylamine, 4-(Sulfonamidomethyl)benzylamine, 2-Chloro-6-fluoro-benzylamine, 3-Chloro-4-fluoro-benzylamine, 2-Chloro-4-fluoro-benzylamine, 3,5-
- 20 Difluoro-benzylamine, 2,4-Difluoro-benzylamine. Pentafluorobenzylamine, 4-Methoxy-2,3,5,6-tetrafluorobenzylamine, 4-(Methyl)benzylamine, Benzylamine, 4-(Ethyl)benzylamine, 4-(Ethoxy)benzylamine, 4-(Isopropyl)benzylamine, 4-(Isobutyl)benzylamine, 4-(Isopropropoxy)benzylamine, 4-
- (Isobutoxy)benzylamine, 4-(Allyl)benzylamine, 4-25 (Allyloxy)benzylamine, 4-(3,3,1,1-Tetrafluoroallyloxy)benzylamine, 4-(Trifluoromethoxy)benzylamine, 4-(2,2,2-trifluoroethoxy)benzylamine, 3,4-Ethylenedioxybenzylamine, 4-Methoxymethyl-2-chlorophenethylamine, 4-(2-Methoxyethyl)phenethylamine, 4-
- (Ethoxymethyl)phenethylamine, 4-(Acetoxyoxymethyl)phenethylamine, 30 3-(Dimethylaminomethyl)phenethylamine, 1-Phenyl-2,2,2trifluoroethylamine, 4-(Trifluoromethoxy)aniline, 4-Methoxyaniline, 4-Ethoxyaniline, 3-Chloro-4-fluoro-aniline, 4-Chloro-2-fluoro-aniline, 4-(Acetoxy)aniline, 4-(Butoxy)aniline, 3-Chloroaniline, 4-

- (Methylthio)aniline, 5-(Aminomethyl)benzofuran, 5-(Methylaminomethyl)benzofuran, 4-(1-Aminoethyl)benzofuran, 5-(2-Aminoethyl)benzofuran, 5-Aminomethyl-2,3-dihydro-benzofuran, 5-Methylaminomethyl-2,3-dihydro-benzofuran, 4-1-Aminoethyl-2,3-dihydro-benzofuran, 5-Aminomethyl-2,3-dihydro-benzofuran, 5
- dihydro-benzofuran, 5-2-Aminoethyl-2,3-dihydro-benzofuran, 5-Aminomethyl-2H-tetrahydrobenzopyran, 5-Methylaminomethyl-2H-tetrahydrobenzopyran, 4-1-Aminoethyl-2H-tetrahydrobenzopyran, 5-Aminomethyl-2H-tetrahydrobenzopyran, 5-Aminomethyl-2H-tetrahydrobenzopyran, 5-Methylaminomethyl-2H-tetrahydrobenzopyran,
- 4-(1-Aminoethyl)-2H-tetrahydrobenzopyran, 5-(2-Aminoethyl)-2H-tetrahydrobenzopyran, 5-Aminomethyl-benzo-1,4-dioxane, 5-Methylaminomethyl-benzo-1,4-dioxane, 4-1-Aminoethyl-benzo-1,4-dioxane, 5-2-Aminoethyl-benzo-1,4-dioxane, 5-Aminomethyl-benzo-1,4-dioxane, 5-Methylaminomethyl-benzo-1,4-dioxane, 4-(1-Aminoethyl)-
- benzo-1,4-dioxane, 5-(2-Aminoethyl)-benzo-1,4-dioxane, 3-Amino-5-methoxy-thiophene, 2-Amino-5-chloro-thiophene, 2-(2-Aminoethyl)thiophene, 2-(3-Aminopropyl)thiophene, 3-(3-Aminopropyl)thiophene, 3-(2-Methylaminoethyl)thiophene, 2-Chloro-3-(2-aminoethyl)-thiophene, 2-Aminoethyl-4-methoxy-thiophene, 2-
- Amino-3-ethyl-thiophene, 2-(Methylaminomethyl)thiophene, 3(Aminomethyl)thiophene, 2-(2-Aminoethyl)-4-methoxy-thiophene, 1(Aminomethyl)tetrazole, 1-(1-Aminoethyl)tetrazole, 1-(3-Aminopropyl)tetrazole, 5-Amino-3-methyl-isoxazole, 3-Aminopyridine, 4-Aminomethylthiazole, 2-(2-Aminoethyl)pyrazine, 2-(1-
- 25 Aminoethyl)imidazole, 2-(Aminomethyl)isoxazole, 3-(2-Aminoethyl)pyrazole, 2-(Aminomethyl)-1,3,4-thiadiazole.

EXAMPLE 305 Methyl 29,30,31,32-tetrahydro-nodulisporate

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To 1.3 mg methyl nodulisporate in 2 mL 1:1 benzene/water at room temperature was added 1 drop Adogen® 464 (Aldrich Chemical Co., Milwaukee, Wisconsin), 10 mg NaHCO3 and 10 mg Na₂S₂O₄. The solution was heated to 80°C for 10 minutes. The reaction was cooled to

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room temperature, extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated under reduced pressure. Purified product was obtained following preparative TLC (1 x 0.5 mm silica gel) using 6:4 EtOAc/hexanes as eluant. The purified product was characterized by ¹H NMR.

EXAMPLE 306

N-(2-Tetrahydrofuranylmethyl)-29,30,31,32-tetrahydro-nodulisporamide

To 40 mg N-(2-tetrahydrofuranylmethyl)-nodulisporamide in 2 mL methanol at room temperature was added 20 mg 10% Pd on carbon. One atmosphere of hydrogen was established and maintained for 2 hours using a balloon. After removal of the catalyst by filtration through Celite using methanol as eluant, the solution was concentrated under reduced pressure and 3 mg pure product was obtained following preparative TLC on silica gel (two 1000 micron plates). The product was characterized by NMR and mass spectrometry (m/z: 767 (M +1)).

EXAMPLE 307

N-Ethyl-N-methyl-29,30,31,32-tetrahydro-nodulisporamide

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To 23 mg of N-ethyl-N-methyl-nodulisporamide in 2 mL methanol at room temperature was added 40 mg 10% Pd on carbon. One atmosphere of hydrogen was established and maintained for 3 hours using a balloon. After removal of the catalyst by filtration through Celite using methanol as eluant, the solution was concentrated under reduced pressure and 9.5 mg of reduced product was obtained following medium pressure liquid chromatography (93/7 methanol/water as eluant). The product was characterized by proton NMR and mass spectrometry (m/z: 723 (M+1)).

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EXAMPLE 308

General Procedure for the Preparation of 29,30,31,32-Tetrahydro-nodulisporic Acid Derivatives

Place 50 mg of a nodulisporamide or nodulisporate analog prepared from the amines listed in Table 6 or the alcohols listed in Table 2 in 4 mL methanol at room temperature. Hydrogenation may be accomplished using 10% Pd on carbon under 1 atmosphere of hydrogen from 15 minutes to 24 hours. The catalyst may be removed by filtration through a pad of Celite using methanol as eluant. Concentration of the solution under reduced pressure followed by purification on silica gel by either flash chromatography, preparative TLC or by reversed-phase liquid chromatography will yield the desired corresponding 29,30,31,32-tetrahydro derivative.

Alternatively, place 50 mg nodulisporic acid in 4 mL methanol at room temperature. Add 1 to 50 mg 10% Pd on carbon and establish an atmosphere of hydrogen using a balloon for 15 minutes to 24 hours. The catalyst may be subsequently removed by filtration through a pad of Celite using methanol as eluant. Concentration of the solution under reduced pressure followed by purification on silica gel by either flash chromatography, preparative TLC or by reversed-phase liquid chromatography will yield the desired corresponding 29,30,31,32-tetrahydro-nodulisporic acid thus obtained may be coupled to the amines in Table 6 or the alcohols listed in Table 2 to form the desired 29,30,31,32-tetrahydro-amide and ester derivatives.

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EXAMPLE 309 29,30-Dihydro-nodulisporic acid

To 1 mg of nodulisporic acid in 1 mL of dichloromethane was added 1.6 mg of Wilkinson's catalyst. The mixture was stirred under a balloon atmosphere of hydrogen overnight (18 h). HPLC separation was obtained with a Magnum 9-ODS reverse phase column and 85:15 methanol:water to 100% methanol gradient. The purified product was isolated upon evaporation of the solvent and characterized by its ¹H NMR.

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EXAMPLE 311

General Procedure for the Preparation of 29,30-Dihydro-Nodulisporic Acid Derivatives

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To a solution of 30 mg of 29,30-dihydro-nodulisporic acid in 3 mL methylene chloride at 0 °C add 0.03 mL triethylamine and 12 mg N-hydroxybenzotriazole followed by 28 mg BOP reagent. Stir the solution for 10 minutes and then add 50 mg of an amine or an alcohol selected from Table 6. Stir overnight at 4 °C and then at room temperature for 2 hours. Pour the solution into 1/1 saturated sodium bicarbonate/brine. Extract the solution with methylene chloride and dry the combined organic layers over sodium sulfate. Remove the solids by filtration and concentrate the solution under reduced pressure. Pure product may be obtained following purification by flash chromatography, preparative TLC or reversed-phase liquid chromatography. Products may be characterized by proton NMR and or mass spectrometry.

EXAMPLE 312

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General Procedure for the Preparation of 31,32-Dihydro-Compound B Derivatives

Place 50 mg of a ester or amide analog prepared from compound B and the amines listed in Table 6 or alcohols listed in Table 2 in 4 mL methanol at room temperature. Hydrogenation of the 31,32-double bond may be accomplished using 10% Pd on carbon under 1 atmosphere of hydrogen from 15 minutes to 24 hours. The catalyst may be removed by filtration through a pad of Celite using methanol as eluant. Concentration of the solution under reduced pressure followed by purification on silica gel by either flash chromatography, preparative TLC or by reversed-phase liquid chromatography will yield the desired 31,32-dihydro-Compound B derivative.

Alternatively, place 50 mg compound B in 4 mL methanol at room temperature. Add 1 to 50 mg 10% Pd on carbon and establish an atmosphere of hydrogen using a balloon for 15 minutes to 24 hours. The catalyst may be subsequently removed by filtration through a pad of Celite using methanol as eluant. Concentration of the solution under reduced pressure followed by purification on silica gel by either flash chromatography, preparative TLC or by reversed-phase liquid chromatography will yield the desired corresponding 31,32-dihydro-compound B. The 31,32-dihydro-compound B thus formed may be coupled to the amines listed in Table 6 and the alcohols listed in Table 2 to form the desired 31,32-dihydro-compound B amides and esters.

EXAMPLE 313 Nodulisporyl azide

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To 1 mg of nodulisporic acid in 0.2 mL chloroform was added 50 μ L triethylamine and 20 μ L of diphenylphosphoryl azide. The reaction mixture was stirred at room temperature for 3h before purification on silica gel (preparative TLC, 1 x 0.5 mm silica gel) using 1:1 EtOAc/hexanes to yield 0.8 mg of pure product which was characterized by 1 H NMR and mass spectrometry.

EXAMPLE 314

29,30-Dihydro-20,30-oxa-nodulisporyl azide

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To 1 mg 29,30-dihydro-20,30-oxa-nodulisporic acid in 0.2 mL chloroform add 0.05 mL triethylamine followed by 0.02 mL diphenylphosphoryl azide. Stir the reaction at room temperature for 3 h before purification by flash chromatography or preparative TLC on silica gel. The product which is obtained may be characterized by proton NMR and mass spectrometry.

EXAMPLE 315

29,30-Dihydro-20,30-oxa-32-descarboxy-32-isocyanato-

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nodulisporic acid

Heat 20 mg of 29,30-dihydro-20,30-oxa-nodulisporyl azide in 8 mL toluene to 90 °C for 2 h. The solvent may be removed by evaporation and the product which is obtained may be characterized by proton NMR and mass spectrometry.

EXAMPLE 316

32-Descarboxy-32-isocyanato-nodulisporic acid

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A solution of 54 mg of nodulisporyl azide in toluene was heated at 90°C for 2 h. The solvent was then evaporated and the isocyanate product was obtained in quantitative yield and was characterized by ¹H NMR and mass spectrometry.

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EXAMPLE 317

32-Descarboxy-32-(1-carbomethoxyamino)-nodulisporic acid

To 1.3 mg of isocyanate of Example 313 in 1 mL of methanol was added 20 microliters of triethylamine. The reaction mixture 20 was heated for 45 min at 75°C and the carbamate product (0.7 mg) was isolated by preparative TLC on silica gel (1 x 0.5 mm) and characterized by ¹H NMR and mass spectrometry.

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EXAMPLE 318

32-Descarboxy-32-(1-(3-benzyl)urea)-nodulisporic acid

To 1 mg of isocyanate of Example 313 in 0.2 mL of toluene was added 40 microliters of benzylamine. The mixture was stirred at 20°C for 20 min and the urea product (0.2 mg) was isolated by 30 preparative TLC (1 x 0.5 mm silica gel, 1:3 hexane:EtOAc) and characterized by its ¹H NMR and MS.

The general procedure of Example 318 was repeated using the appropriate amine to provide urea compounds of Table 7.

Table 7: 32-Descarboxy-32-[UREA]-Nodulisporic Acid Derivatives

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Example	Urea
319	NHC(O)-morpholinyl
320	NHC(O)NHCH ₂ Ph(4-OMe)
321	NHC(O)NHCH(Me) ₂
322	NHC(O)NH(CH ₂) ₅ NH ₂
323	NHC(O)NHCH2CH2OH
333	NHC(O)NHCH2CH2CH2NMe2
334	NHC(O)NHCH2CH2CH2-1-morpholinyl
335	NHC(O)NHCH ₂ -(2-pyridyl)
336	NHC(O)NHCH2CH2-piperazinyl

EXAMPLE 337

General Procedure for the Synthesis of 32-Descarboxy-32-[UREA]- or 32-Descarboxy-32-[CARBAMATE]-Nodulisporic Acid Derivatives

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To 1 mg of isocyanate of Example 313 in 0.2 mL of toluene add 40 mg of an amine selected from Table 6 or alcohol selected from Table 2. Stir the mixture at 20°C from 20 minutes to 24 hours. Pure urea or carbamate product may be isolated by flash chromatography, preparative TLC or reversed-phase liquid chromatography. The purified products may be characterized by proton NMR and mass spectrometry.

EXAMPLE 338

29,30-Dihydro-20,30-oxa-32-descarboxy-32-isocyanato-nodulisporic acid

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Heat a solution of 54 mg of 29,30-dihydro-20,30-oxa-nodulisporyl azide in toluene at 90°C for 2 h. Evaporate the solvent and

the isocyanate product thus obtained may be characterized by ¹H NMR and mass spectrometry.

EXAMPLE 339

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General Procedure for the Synthesis of 29,30-Dihydro-20,30-oxa-32-descarboxy-32-[UREA]- or 29,30-Dihydro-20,30-oxa-32-descarboxy-32-[CARBAMATE]Nodulisporic Acid Derivatives

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To 1 mg of 29,30-dihydro-20,30-oxa-32-descarboxy-32-isocyanato-nodulisporic acid in 0.2 mL of toluene add 40 mg of an amine selected from Table 6 or alcohol selected from Table 2. Stir the mixture at 20°C from 20 minutes to 24 hours. Pure urea or carbamate product may be isolated by flash chromatography, preparative TLC or reversed-phase liquid chromatography. The purified products may be characterized by proton NMR and mass spectrometry.

EXAMPLE 340

31-Hydroxy-20,30-oxa-29,30,31,32-tetrahydro-nodulisporyl azide

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To 1 mg 31-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-nodulisporic acid in 0.2 mL chloroform add 0.05 mL triethylamine followed by 0.02 mL diphenylphosphoryl azide. Stir the reaction at room temperature for 3 h before purification by flash chromatography or preparative TLC on silica gel. The product which is obtained may be characterized by proton NMR and mass spectrometry.

EXAMPLE 341

31-Hydroxy-20,30-oxa-29,30,31,32-tetrahydro-32-descarboxy-32-isocyanato-nodulisporic acid

Heat a solution of 54 mg of 31-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-nodulisporyl azide in toluene at 90°C for 2 h.

Evaporate the solvent and the isocyanate product thus obtained may be characterized by ¹H NMR and mass spectrometry.

EXAMPLE 342

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General Procedure for the Synthesis of 31-Hydroxy-20,30-oxa-32-descarboxy-32-[UREA]-29,30,31,32tetrahydro- or 31-Hydroxy-20,30-oxa-32-descarboxy-32-[CARBAMATE]-29,30,31,32-tetrahydro-nodulisporic acid Derivatives

10 15

To 1 mg of 31-hydroxy-20,30-oxa-29,30,31,32-tetrahydro-32descarboxy-32-isocyanato-nodulisporic acid in 0.2 mL of toluene add 40 mg of an amine selected from Table 6 or alcohol selected from Table 2. Stir the mixture at 20°C from 20 minutes to 24 hours. Pure urea or carbamate product may be isolated by flash chromatography, preparative TLC or reversed-phase liquid chromatography. The purified products may be characterized by proton NMR and mass spectrometry.

EXAMPLE 343 1-Hydroxy-nodulisporic acid

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To 2.8 mg of nodulisporic acid in 0.8 mL of THF at 0°C under argon was added 100 microliters of 2.0 M lithium borohydride in THF. The reaction was quenched with 400 microliters of 2N HCl after 5 min at 0°C and the products were extracted with EtOAc. The extracts were dried over sodium sulfate and concentrated in vacuo. The reside was purified by preparative TLC (1 x 0.5 mm silica gel plate, 95:5:0.5 dichloromethane:methanol:acetic acid) to yield 0.8 mg of isomer A and 0.6 mg of isomer B characterized by their ¹H NMR and MS.

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EXAMPLE 344

1-Hydroxy-nodulisporic acid, methyl ester

To 0.5 mg methyl nodulisporate in 1 mL methanol at 0°C was added 1 mg sodium borohydride. After 10 min at 0°C, the solution WO 96/29073 PCT/US96/03611

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was purified by reversed-phase HPLC without workup using 30:70 to 15:85 (25 minute linear gradient) water/methanol to yield pure product. The product was characterized by ¹H NMR.

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EXAMPLE 345 N-Ethyl-N-methyl-1-hydroxy-nodulisporamide

To 30 mg N-ethyl-N-methyl-nodulisporamide in 2 mL tetrahydrofuran at room temperature was added 1 mL diisobutylaluminum hydride (1.0 M solution in hexanes). After 3 days at room temperature, the reaction was quenched by the addition of acetic acid. The solution was washed with saturated sodium bicarbonate and brine, dried over sodium sulfate and evaporated to dryness. The residue was purified by flash chromatography on silica gel using 1/1 acetone/hexanes as eluant. The purified product was characterized by proton NMR and mass spectrometry (m/z: 723 (M+1)).

EXAMPLE 346 1-Hydroxy-Compound B or C

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To 5 mg of Compound B or C in 2 mL of methanol at 0°C under argon add 5 mg of sodium borohydride. After 10 min at 0°C, extract the products with methylene chloride. Dry the combined extracts over sodium sulfate and concentrate the solution in vacuo. The residual solid may be purified purified by flash chroimatography, preparative TLC or reversed-phase liquid chromatography to yield 1-hydroxy-Compound B or C as a mixture of stereoisomers which may be characterized by proton NMR and mass spectrometry.

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EXAMPLE 347

General Procedure for Synthesis of 1-Hydroxy-Amide and Ester Derivatives of Compounds A, B and C

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To a solution of 30 mg of 1-hydroxy-Compound A, B or C in 3 mL methylene chloride at 0°C add 0.03 mL triethylamine and 12 mg N-hydroxybenzotriazole followed by 28 mg BOP reagent. Stir the solution for 10 minutes and then add 50 mg of an amine selected from Table 6 or an alcohol selected from Table 2. Stir overnight at 4 °C and then at room temperature for 2 hours. Pour the solution into 1/1 saturated sodium bicarbonate/brine. Extract the solution with methylene chloride and dry the combined organic layers over sodium sulfate. Remove the solids by filtration and concentrate the solution under reduced pressure. Pure product may be obtained following purification by flash chromatography, preparative TLC or reversed-phase liquid chromatography. Products may be characterized by proton NMR and/or mass spectrometry.

EXAMPLE 348

1-Hydroxy-1-methyl-nodulisporic acid

To 0.5 mL of 1.4 M methylmagnesium bromide in THF/toluene at 0°C was added 1 mg of nodulisporic acid dissolved in 0.6 mL of THF. After 10 min, the reaction was quenched with 2N HCl and extracted with EtOAc. Preparative TLC (1 x 0.5 mm silica gel plate, 95:5:0.5 dichloromethane:methanol:acetic acid) gave 0.8 mg of product characterized by its ¹H NMR.

EXAMPLE 349

1-Hydroxy-1-methyl-nodulisporic acid, methyl ester

To 1.2 mg of methyl nodulisporate in 1 mL of THF under argon at -78°C was added 0.5 mL of 1.4M methylmagnesium bromide in THF/toluene. The mixture was stirred for 15 min before an aqueous solution of ammonium chloride was added. The mixture was extracted with EtOAc. Preparative TLC (1 x 0.5 mm silica gel plate, 2:3 hexane:EtOAc) gave 1 mg of the titled product characterized by its ¹H NMR.

EXAMPLE 350

1-Hydroxy-1-Alkyl- or 1-Hydroxy-1-Aryl-Compounds A, B or C

To 0.5 mL solution of 1.0 M Grignard reagent selected from Table 8 in 1/1 THF/toluene at 0°C add 1 mg Compound A, B or C dissolved in 0.6 mL THF. After 10 min at 0°C, quench the reaction with 2N HCl and extract with methylene chloride. Dry the combined organic layers over sodium sulfate, filter and concentrate under reduced pressure. Pure product may be obtained following flash chromatography, preparative 10 TLC or reversed-phase liquid chromatography. Purified products may be characterized by proton NMR or mass spectrometry.

Table 8: Grignard Reagents

15 Methyl magnesium bromide
Ethyl magnesium chloride
iso-Propyl magnesium bromide
Phenyl magnesium iodide
Benzyl magnesium bromide

20 Allyl magnesium bromide

O Allyl magnesium bromide
Propargyl magnesium bromide
Magnesium bromide acetilide

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EXAMPLE 351

25 1-Hydroxy-32-descarboxy-32-hydroxymethyl-nodulisporic acid

To 1.2 mg methyl nodulisporate in 1.2 mL tetrahydrofuran at -78°C was added 20 µL 1M lithium aluminum hydride in tetrahydrofuran. The yellow color rapidly disappeared. After 10 minutes, the reaction was quenched at -78°C by dropwise addition of saturated Na₂SO₄. The solution was extracted with ethyl acetate, dried with Na₂SO₄, filtered and concentrated under reduced pressure. Pure product was obtained following preparative TLC (1 x 0.25 mm silica gel

plate) using 85:15 EtOAc/hexanes as eluant. The purified product was characterized by ¹H NMR.

EXAMPLE 352

5 31,32-Dihydro-31,32-dihydroxy-nodulisporic acid and Aldehyde (Compound IV)

To 3 mg of nodulisporic acid was added 1 mL of methanol and 100 microliters of 0.04 M OsO4 in t-butanol stabilized with 1% t-butyl hydroperoxide. After 50 min at room temperature, 400 mg of sodium sulfite in 2 mL of water was then added to the reaction mixture and stirring was continued for another 20 minutes. The mixture was then extracted with EtOAc and the crude products were purified by preparative TLC (1 x 0.5 mm silica gel plate) eluted in 95:5:0.5 dichloromethane:methanol:acetic acid to yield the title compound (1 mg isomer A and 0.6 mg isomer B) and 0.5 mg of aldehyde derived from nodulisporic acid (Compound IV), each characterized by ¹H NMR.

EXAMPLE 353

General Procedure for the Preparation of Ester and Amide Derivatives of 31,32-Dihydro-31,32-dihydroxy-nodulisporic acid

To a solution of 30 mg of 31,32-dihydro-31,32-dihydroxynodulisporic acid in 3 mL methylene chloride at 0 °C add 0.03 mL
triethylamine and 12 mg N-hydroxybenzotriazole followed by 28 mg
BOP reagent. Stir the solution for 10 minutes and then add 50 mg of
amine listed in Table 6 or an alcohol listed in Table 2. Stir the solution
overnight at 4 °C and then pour into 1/1 saturated sodium
bicarbonate/brine, extract with methylene chloride and dry the combined
organic layers over sodium sulfate. Remove the solids by filtration and
concentrate the solution to dryness under reduced pressure. Pure product
may be obtained by flash chromatography or preparative TLC on silica
gel or reversed-phase liquid chromatography. The purified product may
be characterized by proton NMR and mass spectrometry.

EXAMPLE 354 4,20-bis-O-Acetyl-nodulisporic acid

To 1.2 mg of nodulisporic acid was added 300 microliters of acetic anhydride and 100 microliters of pyridine. The reaction mixture was heated at 65°C for 1 h and excess solvent was removed in vacuo. The residual solid was purified by preparative TLC on silica gel eluted with 95:5 dichloromethane:methanol to yield 1.2 mg of the bis-acetate characterized by its ¹H NMR.

EXAMPLE 355

N-Ethyl-N-methyl-20-dimethylaminocarbonyloxy-nodulisporamide

To 30 mg N-ethyl-N-methyl-nodulisporamide in 3 mL methylene chloride at 4 °C was added 60 mg carbonyldiimidazole. After 3 days at 4 °C, 1 mL dimethylamine (25% in water) was added and the solution stirred for an additional 4 days. The solution was poured into brine, extracted with methylene chloride, dried with sodium sulfate and evaporated to dryness. Product was partially purified by flash chromatography on silica gel using 4/6 acetone/hexanes as eluant. Additional purification using medium pressure liquid chromatography (92/8 methanol/water as eluant) yielded 18 mg pure product. The purified product was characterized by proton NMR and mass spectrometry (m/z: 792 (M+1)).

EXAMPLE 356

N-Ethyl-N-methyl-1-desoxo-1-methoximino-nodulisporamide

To a solution of 30 mg N-ethyl-N-methyl-nodulisporamide and 30 mg methoxylamine hydrochloride in 4 mL ethanol was added 0.1 mL pyridine. The solution was heated to reflux for 2 days, cooled to room temperature and concentrated under reduced pressure. The residue was diluted with methylene chloride, washed with brine, dried over sodium

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sulfate and concentrated to dryness. The residue was purified by preparative TLC on silica gel (two 1000 micron plates) using 1/9 methanol/methylene chloride as eluant. The purified products (26 mg), as a mixture of E- and Z-methoximes, were characterized by proton NMR and mass spectrometry (m/z: 732 (M+1 - 1H2O)).

EXAMPLE 357

N-Ethyl-N-methyl-1-desoxo-1-oximino-nodulisporamide

To a solution of 20 mg N-ethyl-N-methyl-nodulisporamide and 20 mg hydroxylamine hydrochloride in 2 mL ethanol at room temperature was added 0.02 mL pyridine. The solution was heated to reflux for 15 hours, cooled to room temperature and diluted with methylene chloride. The solution was washed with brine, dried over sodium sulfate and concentrated to dryness. The residue was purified by preparative TLC on silica gel (two 1000 micron plates) using 1/9 methanol/methylene chloride as eluant to yield 17 mg desired product as a mixture of E- and Z-oxime isomers. The purified products were characterized by proton NMR and mass spectrometry (m/z: 718 (M+1 - 1H2O)).

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EXAMPLE 358

General Procedure for the Preparation of 1-Oximino Derivatives of Compounds A, B and C

To a solution of 20 mg of compound A, B or C and 20 mg hydroxylamine derivative selected from Table 9 in 2 mL ethanol at room temperature, add 0.02 mL pyridine. Heat the solution to reflux for 15 minutes to 24 hours, then cool to room temperature and dilute with methylene chloride. The solution may be washed with brine, the organic layer dried over sodium sulfate and concentrated to under reduced pressure. Pure product may be obtained following purification by flash chromatgraphy or preparative TLC on silica gel or reversed-phase liquid chromatography as a mixture of E- and Z-oxime isomers. The purified products may be characterized by proton NMR and mass spectrometry.

Similarly, amide and ester derivatives of compounds A, B and C, prepared using the amines listed in Table 6 and alcohols in Table 2, may be substituted for compounds A, B and C in the above procedure.

5 Table 9: Oxime Reagents

Hydroxylamine

- O-Methylhydroxylamine
- O-Ethylhydroxylamine
- 10 O-Benzylhydroxylamine
 - O-tert-Butylhydroxylamine
 - O-(Pentafluorobenyzl)hydroxylamine
 - O-Allylhydroxylamine
 - O-Phenylhydroxylamine
- 15 O-iso-Butylhydroxylamine

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- O-(2-Chloro-6-fluoro-benzyl)hydroxylamine
- O-(4-Methoxybenzyl)hydroxylamine

EXAMPLE 359

General Procedure for the Preparation of Hydrazinyl Derivatives of Compounds A, B and C

To a solution of 20 mg of compound A, B or C and 20 mg hydrazine selected from Table 10 in 2 mL ethanol at room temperature, add 0.02 mL pyridine. Heat the solution to reflux for 15 minutes to 24 hours, then cool to room temperature and dilute with methylene chloride. The solution may be washed with brine, the organic layer dried over sodium sulfate and concentrated to under reduced pressure. Pure product may be obtained following purification by flash chromatgraphy or preparative TLC on silica gel or reversed-phase liquid chromatography as a mixture of E- and Z-oxime isomers. The purified products may be characterized by proton NMR and mass spectrometry. Similarly, amide and ester derivatives of compounds A, B and C, prepared using the

amines listed in Table 6 and alcohols in Table 2, may be substituted for compounds A, B and C in the above procedure.

Table 10: Hydrazine Reagents

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Methylhydrazine N,N-Dimethylhydrazine tert-Butylhydrazine 4-Amino-morpholine

10 1-Amino-pyrrolidine

1-Amino-piperidine

Phenylhydrazine

4-(Methyl)phenylhydrazine

Benzylhydrazine

15 Ethyl hydrazinoacetate

2-(Fluoro)phenylhydrazine

1-Amino-4-methyl-piperazine

1-Amino-4-(2-hydroxyethyl)piperazine

2,5-Dichlorophenylhydrazine

20 Methanesulfonyl hydrazide iso-Propylsulfonyl hydrazide Benzenesulfonyl hydrazide

EXAMPLE 360

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N-Ethyl-N-methyl-26-epi-nodulisporamide

To a solution of 5 mg N-ethyl-N-methyl-nodulisporamide in 2 mL acetonitrile was added 1 mL triethylamine. The solution was heated to reflux for 20 hours. The solution was concentrated to dryness under reduced pressure. The residue was purified by flash chromatography on silica gel using 1/9 methanol/methylene chloride to yield the desired product, which was characterized by proton NMR.

WHAT IS CLAIMED IS:

1. A compound having the formula I:

wherein

5

R₁ is **(1)** hydrogen, optionally substituted C1-C10 alkyl, 10 (2) optionally substituted C2-C10 alkenyl, (3) **(4)** optionally substituted C2-C10 alkynyl, optionally substituted C3-C8 cycloalkyl, **(5)** optionally substituted C5-C8 cycloalkenyl (6) where the substitutents on the alkyl, alkenyl, alkynyl, 15 cycloalkyl and cycloalkenyl are 1 to 3 groups independently selected from (i) C₁-C₅ alkyl, (ii) X-C1-C10 alkyl, where X is O or S(O)_m. 20 C3-C8 cycloalkyl, (iii) (iv) hydroxy, (v) halogen,

cyano, carboxy,

25 (viii) NY¹Y², where Y¹ and Y² are independently hydrogen or C₁-C₁₀ alkyl,

(vi)

(vii)

(ix) C1-C10 alkanoylamino, and

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- (x) aroyl amino wherein said aroyl is optionally substituted with 1 to 3 groups independently selected from Rf
- (7) aryl C0-C5 alkyl wherein said aryl is optionally substituted with 1 to 3 groups independently selected from Rf.
- (8) C₁-C₅ perfluoroalkyl
- (9) a 5- or 6-membered heterocycle containing from 1 to 4 heteroatoms independently selected from oxyger, sulfur and nitrogen atoms optionally substituted by 1 to 3 groups independently selected from hydroxy, oxo, C1-C10 alkyl and halogen, and which may be saturated or partly unsaturated,

R2, R3, and R4 are independently ORa, OCO₂Rb, OC(O)NRcRd; or R1+R2 represent =O, =NORa or =N-NRcRd; R5 and R6 are hydrogen; or

R5 and R6 together represent -O-;

R7 is (1) CHO, or

(7)

	R8 is	(1)	hydrogen,
20		(2)	ORa, or
		(3)	NRcRd
	R9 is	(1)	hydrogen, or
		(2)	ORa;
	R10 is	(1)	CN,
25		(2)	$C(O)OR^b$,
		(3)	$C(O)N(OR^b)R^c$
		(4)	C(O)NRcRd,
		(5)	NHC(O)ORb,
		(6)	NHC(O)NRCRd,

CH₂OR^a,

		(8)	CH2OCO2Rb,
		(9)	CH2OC(O)NRCRd,
		(10)	C(O)NRCNRCRd, or
		(11)	C(O)NRcSO2Rb;
5		represents a s	single or a double bond;
	Ra is	(1)	hydrogen,
		(2)	optionally substituted C1-C10 alkyl,
		(3)	optionally substituted C3-C10 alkenyl,
		(4)	optionally substituted C3-C10 alkynyl,
10		(5)	optionally substituted C1-C10 alkanoyl,
		(6)	optionally substituted C3-C10 alkenoyl,
		(7)	optionally substituted C3-C10 alkynoyl,
		(8)	optionally substituted aroyl,
		(9)	optionally substituted aryl,
15		(10)	optionally substituted C3-C7 cycloalkanoyl,
		(11)	optionally substituted C5-C7 cycloalkenoyl,
		(12)	optionally substituted C1-C10 alkylsulfonyl
		(13)	optionally substituted C3-C8 cycloalkyl
		(14)	optionally substituted C5-C8 cycloalkenyl
20		where the	he substituents on the alkyl, alkenyl, alkynyl,
			l, alkenoyl, alkynoyl, aroyl, aryl, cycloalkanoyl,
		cycloall	cenoyl, alkylsulfonyl, cycloalkyl and cycloalkenyl
		are fron	1 1 to 10 groups independently selected from
		hydroxy	, C1-C6 alkoxy, C3-C7 cycloalkyl, aryl C1-C3
25		alkoxy,	NRgRh, CO ₂ Rb, CONR ^c Rd and halogen,
		(15)	C1-C5 perfluoroalkyl,
		(16)	arylsulfonyl optionally substituted with 1 to 3
		groups i	ndependently selected from C1-C5 alkyl, C1-C5
		perfluor	oalkyl, nitro, halogen and cyano,
30		(17)	a 5- or 6-membered heterocycle containing 1 to 2
		heteroat	oms selected from oxygen, sulfur and nitrogen
		optional	ly substituted by 1 to 4 groups independently
		selected	from C1-C5 alkyl, C1-C5 alkenyl, C1-C5

		perfluoroalkyl, amino, C(O)NRcRd, cyano, CO2Rb and
		halogen, and which may be saturated or partly unsaturated;
	Rb is	(1) hydrogen,
	10 15	(2) optionally substituted aryl,
5		(3) optionally substituted C ₁ -C ₁₀ alkyl,
		1 5 010 untolly1,
		1
10		1 Julian Julian State of Clours of Charles o
10		(8) optionally substituted 5- to 10-membered
		heterocycle containing from 1 to 4 heteroatoms
		independently selected from oxygen, sulfur and nitrogen;
		where the substituents on the aryl, alkyl, alkenyl, cycloalkyl,
15		cycloalkenyl, heterocycle, or alkynyl are from 1 to 10 groups
15		independently selected from
		(i) hydroxy,
		(ii) C ₁ -C ₆ alkyl,
		(iii) oxo,
20		(iv) SO ₂ NRgRh,
20		(v) aryl C ₁ -C ₆ alkoxy,
		(vi) hydroxy C ₁ -C ₆ alkyl,
		(vii) C1-C12 alkoxy,
		(viii) hydroxy C ₁ -C ₆ alkoxy,
25		(ix) amino C ₁ -C ₆ alkoxy,
23		(x) cyano,
		(xi) mercapto,
		(xii) C ₁ -C ₆ alkyl-S(O)m,
		(xiii) C3-C7 cycloalkyl optionally substituted
30		with 1 to 4 groups independently selected from Re,
30		(xiv) C5-C7 cycloalkenyl,
		(xv) halogen,
		(xvi) C ₁ -C ₅ alkanoyloxy,
		(xvii) C(O)NRgRh,
		(xviii) CO ₂ Ri,

		(xix)	formyl,
		(xx)	-NRgR ^h ,
		(xxi)	5 to 9-membered heterocycle, which may
	be	e saturated or par	rtially unsaturated, containing from 1 to 4
5			endently selected from oxygen, sulfur and
	ni	trogen, and optic	onally substituted with 1 to 5 groups
	in	dependently sele	ected from Re,
		(xxii)	optionally substituted aryl, wherein the
	ar	yl substituents a	re 1,2-methylenedioxy or 1 to 5 groups
10	in	dependently sele	ected from Re,
		(xxiii)	optionally substituted aryl C1-C3 alkoxy,
	w	herein the aryl s	ubstituents are 1,2-methylenedioxy or 1 to
	5	groups independ	ently selected from Re, and
	_		C ₁ -C ₅ perfluoroalkyl;
15	R ^c and R ^d are	independently se	elected from R ^b ; or
	R ^c and R ^d toge	ther with the N	to which they are attached form a 3- to 10-
	m	embered ring co	ntaining 0 to 2 additional heteroatoms
	se	lected from O, S	(O) _m , and N, optionally substituted with 1
	to	3 groups indepe	ndently selected from Rg, hydroxy, thioxo
20		d oxo;	
	R^e is (1		
) C1-C7 al	kyl,
	(3)	-	erfluoroalkyl,
	(4)	$-S(O)_{m}R$.1 ,
25	(5)) cyano,	
	(6)		
	(7)		
	(8)		
	(9)) RiOCO(0	$CH_2)_V$,
30	(1)	O) optionall	y substituted aryl where the substituents
	are	from 1 to 3 of 1	nalogen, C1-C6 alkyl, C1-C6 alkoxy, or
	hy	droxy,	
	(1)	I) SO ₂ NRg	R ^h , or
	(12	2) amino;	

	Rf is	(1)	C ₁ -C ₄ alkyl,
		(2)	X-C ₁ -C ₄ alkyl, where X is O or $S(O)_m$,
		(3)	C2-C4 alkenyl,
		(4)	C2-C4 alkynyl,
5		(5)	C1-C3-perfluoroalkyl,
		(6)	NY ¹ Y ² , where Y ¹ and Y ² are independently H or
		C1-C5	s alkyl,
		(7)	hydroxy,
		(8)	halogen, and
10		(9)	C ₁ -C ₅ alkanoyl amino,
	Rg and R	^h are inde	pendently
		(1)	hydrogen,
		(2)	C1-C6 alkyl optionally substituted with hydroxy,
		amino	, or CO ₂ R ⁱ
15		(3)	aryl optionally substituted with halogen, 1,2-
		methy)	lenedioxy, C1-C7 alkoxy, C1-C7 alkyl or C1-C3
		perfluc	proalkyl,
		(4)	aryl C1-C6 alkyl, wherein the aryl is optionally
		substit	uted with C1-C3 perfluorolkyl or 1,2-methylenedioxy;
20		(5)	C1-C5 alkoxycarbonyl,
		(6)	C1-C5 alkanoyl,
		(7)	C1-C5 alkanoyl C1-C6 alkyl,
		(9)	aryl C1-C5 alkoxycarbonyl,
		(10)	aminocarbonyl,
25		(11)	C ₁ -C ₅ monoalkylaminocarbonyl
		(12)	C ₁ -C ₅ dialkylaminocarbonyl; or
	Rg and Rh	together	with the N to which they are attached form a 3- to 7-
		membe	red ring containing 0 to 2 additional heteroatoms
		selected	from O, S(O) _m , and N, optionally substituted with 1
30	_ • .	to 3 gro	oups independently selected from Re and oxo;
	R ⁱ is	(1)	hydrogen,
		(2)	C1-C3 perfluoroalkyl,
		(3)	C ₁ -C ₆ alkyl,

		(4)	4 •	11
		(4)		ally substituted aryl Co-C6 alkyl, where the
		-		are from 1 to 3 groups independently
		selected	from ha	logen, C1-C6 alkyl, C1-C6 alkoxy, and
		hydrox	y;	
5	m is	0 to 2;	and	
	v is	0 to 3 ;	or	
	a pharmac	eutically a	cceptable	e salt thereof; and
	excluding:	nodulispo	ric acid, 2	29,30-dihydro-20,30-oxa-nodulisporic acid
		and 31-	hydroxy-	20,30-oxa-29,30,31,32-tetrahydro-
10		nodulis	poric acid	l.
		2.	A com	oound of Claim 1
	wherein			
	R ₁ is	(1)	hydrog	en.
15	•	(2)		illy substituted C1-C6 alkyl,
		(3)	_	illy substituted C2-C6 alkenyl,
		(4)		illy substituted C2-C6 alkynyl,
		(5)		lly substituted C5-C6 cycloalkyl,
		(6)		lly substituted C5-C6 cycloalkenyl
20		, ,	=	utents on the alkyl, alkenyl, alkynyl,
				cloalkenyl are 1 to 3 groups independently
		selected		general and a group morponically
			(i)	C ₁ -C ₃ alkyl,
			(ii)	X-C1-C6 alkyl, where X is O or $S(O)_m$,
25			(iii)	C5-C6 cycloalkyl,
			(iv)	hydroxy,
			(v)	halogen,
			(vi)	cyano,
			(vii)	carboxy, and
30			(viii)	$NY^{1}Y^{2}$, where Y^{1} and Y^{2} are
			indepen	dently hydrogen or C1-C6 alkyl,
		(7)	_	-C3 alkyl wherein said aryl is optionally
		substitu		to 3 groups independently selected from
		Rf,		

		(8)	C1 C2 porflyons all and
		(9)	C1-C3 perfluoroalkyl,
		- ·	a 5- or 6-membered heterocycle containing from 1
		ond ni	eteroatoms independently selected from oxygen, sulfur
5		indend	trogen atoms optionally substituted by 1 to 3 groups
,		halaaa	endently selected from hydroxy, oxo, C1-C6 alkyl and
	R8 is	natoge	en, and which may be saturated or partly unsaturated,
	1/9 12	(1)	hydrogen,
		(2)	OH, or
10	R9 is	(3)	NH ₂ ;
10	K9 18	(1)	hydrogen or
	Diois	(2)	OH;
	R_{10} is	(1)	C(O)ORb,
		(2)	C(O)N(ORb)Rc,
1.5		(3)	C(O)NRcRd,
15		(4)	NHC(O)ORb,
		(5)	NHC(O)NRCRd,
		(6)	CH ₂ ORa,
		(7)	CH2OCO2Rb,
20		(8)	CH2OC(O)NRcRd,
20		(9)	C(O)NRCNRCRd, or
	7 0.	(10)	C(O)NRcSO2Rb;
	Ra is	(1)	hydrogen,
		(2)	optionally substituted C ₁ -C ₆ alkyl,
0.5		(3)	optionally substituted C3-C6 alkenyl,
25		(4)	optionally substituted C3-C6 alkynyl,
		(5)	optionally substituted C1-C6 alkanoyl,
		(6)	optionally substituted C3-C6 alkenoyl,
		(7)	optionally substituted C3-C6 alkynoyl,
		(8)	optionally substituted aroyl,
30		(9)	optionally substituted aryl,
		(10)	optionally substituted C5-C6 cycloalkanoyl,
		(11)	optionally substituted C5-C6 cycloalkenoyl,
		(12)	optionally substituted C1-C6 alkylsulfonyl
		(13)	optionally substituted C5-C6 cycloalkyl

		alkanoyl	e substitu , alkenoy	ly substituted C5-C6 cycloalkenyl ents on the alkyl, alkenyl, alkynyl, l, alkynoyl, aroyl, aryl, cycloalkanoyl, cylsulfonyl, cycloalkyl and cycloalkenyl
5		are from hydroxy,	1 to 10 g C ₁ -C ₄ a	roups independently selected from lkoxy, C5-C6 cycloalkyl, aryl C1-C3 CO2Rb, CONRCRd and halogen,
		(15)	C ₁ -C ₃ p	erfluoroalkyl,
		(16)	arylsulfo	onyl optionally substituted with 1 to 3
10		groups ir	ndepende	ntly selected from C1-C3 alkyl, C1-C3
		perfluoro	oalkyl, ha	logen and cyano,
		(17)	a 5- or 6	-membered heterocycle containing 1 to 4
		heteroato	oms selec	ted from oxygen, sulfur and nitrogen
		optionall	y substitu	ated by 1 to 4 groups independently
15				C3 alkyl, C1-C3 alkenyl, C1-C3
		perfluoro	oalkyl, am	nino, C(O)NR ^c R ^d , cyano, CO ₂ R ^b and
		halogen,	and whic	h may be saturated or partly unsaturated;
	Rb is	(1)	hydroge	n,
		(2)	-	ly substituted aryl,
20		(3)	_	ly substituted C ₁ -C ₇ alkyl,
		(4)	_	ly substituted C3-C7 alkenyl,
		(5)		ly substituted C3-C7 alkynyl,
		(6)		ly substituted C5-C7 cycloalkyl,
		(7)	-	ly substituted C5-C7 cycloalkenyl, or
25		(8)		ly substituted 5- to 10-membered
		•		ning from 1 to 4 heteroatoms
				ected from oxygen, sulfur and nitrogen;
				ents on the aryl, alkyl, alkenyl, cycloalkyl,
				rocycle, or alkynyl are from 1 to 10 groups
30		independ	-	ected from
			(i)	hydroxy,
			(ii)	C ₁ -C ₃ alkyl,
			(iii)	oxo,
			(iv)	SO ₂ NRgR ^h ,

			•
		(v)	aryl C ₁ -C ₃ alkoxy,
		(vi)	hydroxy C1-C3 alkyl,
		(vii)	C ₁ -C ₇ alkoxy,
		(viii)	hydroxy C1-C3 alkoxy,
5		(ix)	amino C ₁ -C ₃ alkoxy,
		(x)	cyano,
		(xi)	C ₁ -C ₃ perfluoroalkyl,
		(xii)	C ₁ -C ₃ alkyl-S(O)m,
		(xiii)	C5-C6 cycloalkyl optionally substituted
10		with 1	to 4 groups independently selected from Re,
		(xiv)	C5-C6 cycloalkenyl,
		(xv)	
		(xvi)	
		(xvii)	•
15		(xviii)	CO ₂ Ri,
		(xix)	optionally substituted aryl C1-C3 alkoxy,
		wherein the aryl	substituents are 1,2-methylenedioxy or 1 to
		5 groups indepen	ndently selected from Re,
		(xx)	
20		(xxi)	5- to 6-membered heterocycle, which
		may be saturated	or partially unsaturated, containing from 1
		to 4 heteroatoms	independently selected from oxygen, sulfur
		and nitrogen, and	d optionally substituted with 1 to 5 groups
		independently se	lected from Re, and
25		(xxii)	optionally substituted aryl, wherein the
		aryl substituents	are 1,2-methylenedioxy or 1 to 5 groups
		independently se	lected from Re;
	Re is	(1) haloger	
		(2) C_1-C_3	
30		(3) C_1-C_3	perfluoroalkyl,
		$(4) -S(O)_{m}$	R ⁱ ,
		(5) cyano,	
		(6) amino,	
		(7) $R^{i}O(CH)$	f _{2)v} -,

		(8)	R ⁱ CO ₂ (CH ₂) _v -,
		(9)	$R^{i}OCO(CH_2)_{V}$
		(10)	optionally substituted aryl where the substituents
		are from	1 to 3 of halogen, C1-C3 alkyl, C1-C3 alkoxy, or
5		hydroxy	
		(11)	SO ₂ NRgR ^h ;
	Rf is	(1)	methyl,
		(2)	X-C ₁ -C ₂ alkyl, where X is O or S(O) _m ,
		(3)	halogen,
10		(4)	acetylamino,
		(5)	trifluoromethyl,
		(6)	NY ¹ Y ² , where Y ¹ and Y ² are independently H or
		methyl, a	
		(7)	hydroxy;
15	Rg and Rh a	are indepe	ndently
		(1)	hydrogen,
		(2)	C1-C6 alkyl optionally substituted with hydroxy,
		amino, or	
		(3)	aryl optionally substituted with halogen, 1,2-
20		methylen	edioxy, C1-C7 alkoxy, C1-C7 alkyl or C1-C3
		perfluoro	alkyl,
		(4)	aryl C1-C6 alkyl, wherein the aryl is optionally
		substitute	ed with C1-C3 perfluorolkyl or 1,2-methylenedioxy;
		(5)	C ₁ -C ₅ alkoxycarbonyl,
25			C ₁ -C ₅ alkanoyl,
		(7)	C ₁ -C ₅ alkanoyl C ₁ -C ₆ alkyl,
		(9)	aryl C1-C5 alkoxycarbonyl,
		(10)	aminocarbonyl,
		(11)	C ₁ -C ₅ monoalkylaminocarbonyl
30		(12)	C ₁ -C ₅ dialkylaminocarbonyl; or
	Rg and Rh to	agether wi	th the N to subject there are attacked from 5

Rg and Rh together with the N to which they are attached form a 5- to 6membered ring containing 0 to 2 additional heteroatoms selected from O, S(O)_m, and N, optionally substituted with 1 to 3 groups independently selected from Re and oxo;

	ni.	(4)			
	R ⁱ is	(1)	hydrogen,		
		(2)	C ₁ -C ₃ perfluoroalkyl,		
		(3)	C ₁ -C ₄ alkyl,		
		(4)	optionally substituted aryl Co-C4 alkyl, where the		
5		aryl su	ubstituents are from 1 to 3 groups independently		
		selecte	ed from halogen, C1-C4 alkyl, C1-C4 alkoxy, and		
		hydrox			
	all other v	ariables a	re as defined in Claim 1.		
10		3.	A compound of Claim 1		
	wherein	J.	A compound of Claim 1		
	R ₁ is	(1)	hydrogen,		
	1	(2)	optionally substituted C1-C3 alkyl,		
		(3)	optionally substituted C ₂ -C ₃ alkenyl,		
15		(4)			
			optionally substituted C2-C3 alkynyl,		
		1 +0 2 0	he substitutents on the alkyl, alkenyl, and alkynyl are		
		1 to 5 g	roups independently selected from		
			(i) methyl, (ii) X-methyl, where X is O or S(O) _m and		
20					
20		(5)			
			aryl C ₀ -C ₁ alkyl wherein said aryl is optionally		
		Rf,	ted with 1 to 3 groups independently selected from		
		(6)	trifly one action!		
25	R ₈ is	(1)	trifluoromethyl hydrogen,		
	110 15	Ī			
		(2) (3)	OH, or NH2		
	R9 is	(1)	hydrogen, or		
	119 10	(2)			
30	R ₁₀ is	(1)	OH; C(O)OR ^b ,		
	110 15	(2)	$C(O)N(OR^b)R^c$		
		(3)	C(O)N(OR ^o)R ^c , C(O)NR ^c R ^d ,		
		(4)	NHC(O)ORb,		
			· ·		
		(5)	NHC(O)NRcRd,		

		(6)	CH ₂ OR ^a ,
		(7)	CH2OCO2Rb,
		(8)	CH2OC(O)NRCRd,
		(9)	C(O)NRCNRCRd, or
5		(10)	C(O)NRcSO2Rb;
	Ra is	(1)	hydrogen,
		(2)	optionally substituted C1-C4 alkyl,
		(3)	optionally substituted C3-C4 alkenyl,
		(4)	optionally substituted C3-C4 alkynyl,
10		(5)	optionally substituted C1-C4 alkanoyl,
		(6)	optionally substituted aroyl,
		(7)	optionally substituted C5-C6 cycloalkanoyl,
		(8)	optionally substituted C5-C6 cycloalkenoyl,
		(9)	optionally substituted C1-C3 alkylsulfonyl
15		where th	ne substituents on the alkyl, alkenyl, alkynyl,
		alkanoyl	l, aroyl, cycloalkanoyl, cycloalkenoyl, and
		alkylsuli	fonyl, are from 1 to 5 groups independently selected
		from hy	droxy, C1-C2 alkoxy, aryl C1-C3 alkoxy, NRgRh,
		CO ₂ R ^b ,	CONRCRd and halogen,
20		(10)	trifluoromethyl,
		(11)	arylsulfonyl optionally substituted with 1 to 3
		groups in	ndependently selected from methyl, trifluoromethyl
		and halo	gen,
		(12)	a 5- or 6-membered heterocycle containing 1 to 4
25		heteroate	oms selected from oxygen, sulfur and nitrogen
			ly substituted by 1 to 4 groups independently
		selected	from methyl, trifluoromethyl, C(O)NRcRd, CO2Rb
		and halo	gen, and which may be saturated or partly
		unsatura	ted;
30	Rb is	(1)	hydrogen,
		(2)	optionally substituted aryl,
		(3)	optionally substituted C1-C6 alkyl,
		(4)	optionally substituted C3-C6 alkenyl,
		(5)	optionally substituted C3-C6 alkynyl,

	(6) option	ally substituted C5-C6 cycloalkyl,				
	(7) option:	ally substituted C5-C6 cycloalkenyl, or				
	(8) options	ally substituted 5- to 6-membered				
	heterocycle cont	aining from 1 to 4 heteroatoms				
5	independently se	elected from oxygen, sulfur and nitrogen.				
	independently selected from oxygen, sulfur and nitrogen; where the substituents on the aryl, alkyl, alkenyl, cycloalkyl,					
	cycloalkenyl he	terocycle, or alkynyl are from 1 to 10 groups				
	independently se	elected from				
	(i)	hydroxy,				
10	(ii)	C ₁ -C ₃ alkyl,				
	(iii)	oxo,				
	(iv)	SO ₂ NRgR ^h ,				
	(v)	aryl C ₁ -C ₃ alkoxy,				
	(vi)	hydroxy C1-C4 alkyl,				
15	(vii)	C1-C4 alkoxy,				
	(viii)	hydroxy C1-C4 alkoxy,				
	(ix)	amino C ₁ -C ₄ alkoxy,				
	(x)	cyano,				
	(xi)	C_1 - C_4 alkyl- $S(O)m$,				
20	(xii)	C5-C6 cycloalkyl optionally substituted				
	with 1 to	o 4 groups independently selected from Re,				
	(xiii)	C5-C6 cycloalkenyl,				
	(xiv)	halogen,				
	(xv)	C ₁ -C ₃ alkanoyloxy,				
25	(xvi)	C(O)NRgRh,				
	(xvii)	CO ₂ R ⁱ ,				
	(xvii)	-NRgRh,				
	(xix)	5- to 6-membered heterocycle, which				
	may be saturated of	or partially unsaturated, containing from 1				
30	to 4 heteroatoms is	ndependently selected from oxygen, sulfur				
	and nitrogen, and optionally substituted with 1 to 5 groups					
	independently sele	ected from Re,				

			(xx)	optionally substituted aryl, wherein the	
			ostituents are 1,2-methylenedioxy or 1 to 5 groups		
		independ	idently selected from Re,		
			(xxi)	optionally substituted aryl C1-C3 alkoxy,	
5			s independently selected from Re, and		
		5 groups			
	5 0.	443		C ₁ -C ₃ perfluoroalkyl;	
	Re is	(1)	halogen,		
		(2)	C ₁ -C ₃ al		
10			C ₁ -C ₃ perfluoroalkyl,		
		(4)	$-S(O)_{m}R$	1,	
		(5)	cyano,		
		(6)	RiO(CH2		
		(7)	RiCO ₂ (C	— · · ·	
15		(8)	$R^{i}OCO(CH_{2})_{V}$		
		(9)		y substituted aryl where the substituents	
		are from	1 to 3 of 1	nalogen, C1-C3 alkyl, C1-C3 alkoxy, or	
		hydroxy,			
		(10)	SO ₂ NRg	R ^h , or	
20		(11)	amino;		
	Rf is	(1)	methyl,		
		(2)	X-C ₁ -C ₂	alkyl, where X is O or $S(O)_m$,	
		(3)	trifluoron	nethyl,	
		(4)	NY^1Y^2 ,	where Y ¹ and Y ² are independently H or	
25		methyl,	-		
		(5)	hydroxy,		
		(6)	halogen,	and	
		(7)	acetylami	no,	
	Rg and Rh a	re indepe	ndently		
30		(1)	hydrogen	•	
		(2)	C1-C6 all	cyl optionally substituted with hydroxy,	
		amino, or	_	•	

5

- (3) aryl optionally substituted with halogen, 1,2-methylenedioxy, C1-C7 alkoxy, C1-C7 alkyl or C1-C3 perfluoroalkyl,
- (4) aryl C₁-C₆ alkyl, wherein the aryl is optionally substituted with C₁-C₃ perfluorolkyl or 1,2-methylenedioxy;
- (5) C₁-C₅ alkoxycarbonyl,
- (6) C₁-C₅ alkanoyl,
- (7) C₁-C₅ alkanoyl C₁-C₆ alkyl,
- (9) aryl C1-C5 alkoxycarbonyl,
- 10 (10) aminocarbonyl,
 - (11) C₁-C₅ monoalkylaminocarbonyl
 - (12) C₁-C₅ dialkylaminocarbonyl; or

Rg and Rh together with the N to which they are attached form a 5- to 6membered ring containing 0 to 2 additional heteroatoms selected from O, S(O)_m, and N, optionally substituted with 1 to 3 groups independently selected from Re and oxo;

Ri is

- (1) hydrogen,
- (2) C₁-C₃ perfluoroalkyl,
- (3) C₁-C₄ alkyl,

20 (4) optionally substituted aryl C0-C6 alkyl, where the aryl substituents are from 1 to 3 groups independently selected from halogen, C1-C6 alkyl, C1-C6 alkoxy, and hydroxy

all other variables are as defined in Claim 1.

25

- 4. A compound of Claim 1 wherein R7 is CHO.
- 5. A compound of Claim 1 wherein

R7 is the fragment

30

- R₁₀ is
- (1)
- C(O)ORb.

			C(O)N	(OR ^b)R ^c , RcRd, RcNRcRd _{, or}	
				RCSO2Rb	
5	R8, R9, Rb			defined in Claim 1.	
	,				
		6.	A comp	oound of Claim 5 wherein	
	R ₁₀ is	C(O)OI	ζb;		
	Rb is	(1)	optiona	lly substituted aryl,	
10		(2)	optiona	lly substituted C1-C6 alkyl,	
		(3)	optiona	lly substituted C3-C6 alkenyl,	
		(4)	optiona	lly substituted C3-C6 alkynyl,	
		(5)	optiona	lly substituted C3-C6 cycloalkyl, or	
		(6)	optiona	lly substituted 5 to 6-membered heterocycle	
15		containi	containing from 1 to 4 heteroatoms independently selected		
				fur and nitrogen;	
		where th	ne substiti	uents on the aryl, alkyl, alkenyl, cycloalkyl,	
				kynyl are from 1 to 10 groups	
		indepen	dently sel	ected from	
20			(i)	hydroxy,	
			(ii)	C ₁ -C ₃ alkyl,	
			(iii)	oxo,	
			(iv)	SO ₂ NRgR ^h ,	
			(v)	aryl C ₁ -C ₃ alkoxy,	
25			(vi)	hydroxy C ₁ -C ₄ alkyl,	
			(vii)	C ₁ -C ₄ alkoxy,	
			(viii)	hydroxy C ₁ -C ₄ alkoxy,	
			(ix)	amino C1-C4 alkoxy,	
			(x)	cyano,	
30			(xi)	C ₁ -C ₄ alkyl-S(O)m,	
	•		(xii)	C5-C6 cycloalkyl optionally substituted	
			with 1 to	4 groups independently selected from Re,	
			(xiii)	C5-C6 cycloalkenyl,	
			(xiv)	halogen,	

			(xv)	C1-C3 alkanoyloxy,		
			(xvi)	C(O)NRgRh,		
			(xvii)	CO ₂ Ri,		
			(xvii)	-NRgRh,		
5			(xix)	5 to 6-membered heterocycle, which may		
		be satur	ated or pa	rtially unsaturated, containing from 1 to 4		
		heteroat	oms inder	pendently selected from oxygen, sulfur and		
		nitrogen	n, and opti	onally substituted with 1 to 5 groups		
		indepen	dently sele	ected from Re,		
10			(xx)	optionally substituted aryl, wherein the		
		aryl sub	stituents a	re 1,2-methylenedioxy or 1 to 5 groups		
				ected from Re,		
			(xxi)	optionally substituted aryl C1-C3 alkoxy,		
		wherein	the aryl s	ubstituents are 1,2-methylenedioxy or 1 to		
4 groups independently selected from Re, and			lently selected from Re, and			
			(xxii)	C ₁ -C ₃ perfluoroalkyl;		
	Re is	(1)	halogen,			
		(2)	C1-C7 a	lkyl,		
		(3)	C ₁ -C ₃ p	erfluoroalkyl		
20		(4)	nitro,			
		(6)	RiO(CH			
		(7)	RiOC(O)			
		(8)	SO ₂ NR8	gRh,		
	v is	0;				
25	Rg and Rh	Rg and Rh are independently				
		(1)	hydroger	1,		
		(2)	C ₁ -C ₆ al	kyl optionally substituted with hydroxy or		
		CO ₂ Rb,				
		(3)	aryl optic	onally substituted with halogen, 1,2-		
30		methyler	nedioxy, C	1-C7 alkyl or C1-C3 perfluoroalkyl,		
	_	(4)		kanoyl, or		
	Rg and Rh	together w	ith the N	to which they are attached form a 3- to 7-		
		membere	ed ring cor	ntaining 0 to 2 additional heteroatoms		

selected from O, S(O)_m, and N, optionally substituted with 1 to 3 groups independently selected from R^e and oxo;

Ri is

- (1) hydrogen, or
- (2) C_1 - C_6 alkyl;

5 m is

10

0 to 2; and

all other variables are as defined in Claim 5.

7. A compound of Claim 5 wherein

R¹⁰ is

(1) $C(O)N(OR^b)R^c$,

(

- (2) C(O)NRcRd
- (3) C(O)NRCNRCRd, or
- (4) $C(O)NR^{c}SO_{2}R^{i}$;

Rb, Rc, Rd and Ri are as defined in Claim 5.

- 8. A compound of Claim 3 wherein R¹⁰ is C(O)NR^cR^d; and R_c and R^d are as defined in Claim 3.
- 9. A compound of Claim 5 wherein

20 R₁₀ is

30

 $C(O)NR^{c}R^{d};$

Rb is

- (1) hydrogen,
- (2) optionally substituted aryl,
- (3) optionally substituted C1-C6 alkyl,
- (4) optionally substituted C3-C6 alkenyl,
- 25 (5) optionally substituted C3-C6 alkynyl,
 - (6) optionally substituted C3-C6 cycloalkyl,
 - (7) optionally substituted C5-C6 cycloalkenyl, or
 - (8) optionally substituted 5 to 6-membered heterocycle containing from 1 to 4 heteroatoms independently selected from oxygen, sulfur and nitrogen;

where the substituents on the aryl, alkyl, alkenyl, cycloalkyl, cycloalkenyl, heterocycle, or alkynyl are from 1 to 10 groups independently selected from

(i) hydroxy,

	(ii)	C ₁ -C ₃ alkyl,		
	(iii)	oxo,		
	(iv)	SO ₂ NRgRh,		
	(v)	arylC1-C3 alkyl,		
5	(vi)	hydroxy C ₁ -C ₄ alkyl,		
	(vii)	C ₁ -C ₁₂ alkoxy,		
	(viii)	hydroxy C ₁ -C ₄ alkoxy,		
	(ix)	amino C ₁ -C ₄ alkoxy,		
	(x)	cyano,		
10	(xi)	C ₁ -C ₃ perfluoroalkyl,		
	(xii)	C_1 - C_4 alkyl- $S(O)_m$,		
	(xiii)	C5-C6 cycloalkyl optionally substituted		
	with 1 to 4 groups selected from Re,			
	(xiv)	C5-C6 cycloalkenyl,		
15	(xv)	halogen,		
	(xvi)	C(O)NRgRh,		
	(xvii)	CO ₂ R ⁱ ,		
	(xviii)	-NRgRh,		
	(xix)	5 to 9-membered heterocycle containing		
20	from 1 to 4 heteroatoms independently selected from			
	oxygen, sulfur and nitrogen, and optionally substituted with			
	1 to 3 groups independently selected from Re,			
	(xx) optionally substituted aryl, wherein the			
	aryl substituents are 1,2-methylenedioxy or 1 to 5 groups			
25	independently selected from Reand			
	(xxi)	optionally substituted aryl C1-C3 alkoxy,		
	wherein the aryl su	abstituents are 1,2-methylenedioxy or 1 to		
	5 groups independ	ently selected from Re;		
	R ^c and R ^d are independently se	elected from Rb; or		
30	R ^c and R ^d together with the N	to which they are attached form a 3- to 10-		
membered ring containing 0 to 2 additional heteratoms				
	selected from O, S(O)m, and N, optionally substituted with			
		ndently selected from Rg, hydroxy, thioxo		
	and oxo;			

	Re is	(1) (2)	halogen, C1-C3 alkyl,
		(3)	C1-C3 perfluoroalkyl,
		(4)	RiO(CH ₂) _v -,
5		(5)	RjiCO ₂ (CH ₂) _v -,
		(6)	$R^{i}OCO(CH_2)_{v}$
		(7)	SO ₂ NRgRh;
		(8)	amino
	v is	0;	
10	Rg and Rh	Rh are independently	
		(1)	hydrogen,
		(2)	C1-C6 alkyl optionally substituted with hydroxy,
		amino, or CO ₂ Ri,	
		(3)	aryl optionally substituted with halogen, 1,2-
15			nedioxy, C ₁ -C ₇ alkoxy, C ₁ -C ₇ alkyl or C ₁ -C ₃
perfluoroalkyl,			oalkyl,
		(4)	aryl C ₁ -C ₆ alkyl, wherein the aryl is optionally
		substitute	ed with C1-C3 perfluoroalkyl or 1,2-
		methyler	nedioxy,
20		(5)	C ₁ -C ₅ alkoxycarbonyl,
		(6)	C1-C5 alkanoyl,
		(7)	aryl C ₁ -C ₅ alkoxycarbonyl,
		(8)	aminocarbonyl, or
	Rg and Rh	together w	ith the N to which they are attached form a 5- to 6-
25			ed ring containing 0 to 2 additional heteroatoms
	selected from O, S(O)m, and N, optionally substituted with 1		
		to 3 grou	ps independently selected from Re and oxo;
	R ⁱ is	(1)	hydrogen or
		(2)	optionally substituted Co-C6 alkyl wherein the
30		substitue	nts are aryl or substituted aryl, and the aryl
		substitue	nts are from 1 to 3 groups independently selected
			ogen, C1-C6 alkyl, C1-C6 alkoxy, and hydroxy; and
	all other var	iables are	as defined in Claim 5.

10. A compound of Claim 1 wherein R7 is the fragment

CH2ORa, NHC(O)ORb or NHC(O)NRcRd; **R10** is R8, R9, Ra, Rb, Rc, Rd and ____ are as defined in Claim 1.

> 11. A compound of Claim 1 wherein R7 is the fragment

10 **R10** is CO₂H; and R8, R9 and ____ are as defined in Claim 1.

12. A compound of the formula

wherein Rb is selected from the group consisting of: 15

CH3, CH2CH3, CH2CH2OH, CH2CH2N(CH(CH3)2)2, CH2CH2N(CH3)2, CH2CH(OH)CH2N(CH(CH3)2)2,

CH2CH2OCH2CH2OH, CH2Ph(4-NO2), CH2Ph(3-NO2), CH2CF3, 20 CH2CH2CH2C(=O)CH3, CH2CH2CH2Ph, CH2CH2C(CH32CH3, CH(CF3)2, CH2Ph(2-CF3).

13. A compound of the formula

OH

OH

OH

OH

5

wherein R^X is selected from the group consisting of:
H, CH3, CH2CH3, C(CH3)3, CH2CH2CH3, CH2CH2OH,
CH(CO2CH3)CH2OH, CH2CO2CH3, CH2CH(OCH2CH3)2,
CH2CH2OCH2CH2OH, CH(CH3)(CH2)3C(CH3)2OH, (CH2)3OH,
(CH2)4OH, (CH2)5OH, CH(CH2OH)CH2CH3, NHC(CH3)3, CH2CN,
(CH2)6OH, CH2CH(OH)CH3, CH(CH2OH)CH2CH2CH3,
CH2CH2SCH3, CH2CH2SCH2CH3, CH2CONH2,
CH(CH3)(CH2OH)2, CH2CH2NHCH2CH2OH,
CH(CH2OH)(CH2)3CH3, CH(CH2OCH3)CH3, (CH2)2SH,
(CH2)4NH2, CH2CH2SO2CH3, CH2CH2S(O)CH3,
CH(CH(CH3)2)CH2OH, (CH2)3NH2, (CH2)3N(CH2CH3)2,

CH₂)4NH₂, CH₂CH₂SO₂CH₃, CH₂CH₂S(O)CH₃, CH(CH(CH₃)₂)CH₂OH, (CH₂)₃NH₂, (CH₂)₃N(CH₂CH₃)₂, (CH₂)₃N(CH₃)₂, OCH₂CH₃, CH₂CH(OH)CH₂OH, OCH₃, CH₂CH₂OCH₃, CH₂CH₂NHC(O)CH₃, C(CH₃)₂CH₂OH, c-C₃H₅, c-C₆H₁, (CH₂)₃OCH₂CH₃, CH₂CH=CH₂, C(CH₂CH₃)(CH₂OH)₂,

20 CH2C≡CH, CH2CO2CH2CH3, CH2CH2F, (CH2)3O(CH2)11CH3, CH2CH2N(CH3)2, CH2CH2OCH2CH2NH2, CH2CF3, NHCH2CO2CH2CH3, CH(CH3)CO2CH3, C(CH3)2CH2C(O)CH3, CH(CO2CH2CH3)2, CH2CH3, CH(CH2CH2CH3)CO2CH3, CH2CH2CH2CH3)2, CH2CH2CH3, C(CH3)2C≡CH, (CH2)4CH3, CH(CH2CH2CH3)2,

5

(CH₂)₅CH₃,CH₂CH₂CO₂H, CH(CH(CH₃)₂)CO₂CH₃, OCH₂CO₂H, CH(CH(CH₃)₂)CH₂OH, CH(CH(CH₃)₂)CH₂OH, CH(CH₃)CH₂OH, CH(CH₃)₂, C(CH₃)₃, (CH₂)CH(CH₃)₂, CH(CH₃)CH₂CH₃, CH₂CH(CH₃)OH, (CH₂)₃CH₃, (CH₂)₂OCH₂CH₃, 1-adamantyl, (CH₂)₈CH₃, CH(CH₃)CH(CH₃)₂, (CH₂)₃NHCH₃, (CH₂)₂N(CH₂CH₃)₂,

$$-CH_{2}CH_{2}-N \bigcirc O -CH_{2}CH_{2}-N \bigcirc N -CH_{2}-N -CH_$$

15

14. A compound of the formula

wherein NR^xRy is selected from the group consisting of: N(CH₃)CH₂C≡N, N(CH₃)CH₂CH₃, N(CH₃)CH₂CH₃, N(CH₃)CH₂CH₂CH₃, N(CH₃)CH₂CH₂CH₃,

- 5 N(CH₂CH₃)CH₂CH₂OCH₃, N(CH₃)CH₂CH₂OCH₃, N(CH₂CH₃)CH₂CH₂CH₃, N(CH₂CH=CH₂)₂, N(CH₃)CH₂CH₂OH, N(CH₂CH(CH₃)OH)₂, N(CH₂CH₃)₂, N(CH₂CH₂OH)₂, N(CH₂CH₃)CH(CH₃)₂, N(CH₂CH₂CH₃)₂, N(CH₂CH₂CH₂CH₃)₂, N(CH₃)₂, N(CH₂CH₂CH₃)₂,
- N((CH₂)₂CH₃)CH₂CH₂OH, N(CH₃)CH₂C≡CH, N((CH₂)₈CH₃)₂, N((CH₂)₇CH₃)₂, N(CH₃)(CH₂)₂NHCH₃, N(CH₃)(CH₂)₃NH₂, NHCH(CH₂OH)CH₂Ph, NHPh(2-OH,4-CH₃), NHCH₂Ph(4-NH₂), NHPh(4-Cl), NHPh(4-CH₂CH₂OH), NHPh(2-CH₂CH₂OH), NHCH₂CH₂Ph, NHPh(2-CH₂OH), NHPh(3-N(CH₃)₂, NHPh(4-CH₂CH₂OH), NHPh(4-CH₂CH₂OH), NHPh(4-CH₃CH₂OH)
- SO₂NH₂), NHNHPh, NHPh(2-CONH₂), NHCH₂CH₂Ph(4-OH), NHCH₂CH₂Ph(4-SO₂NH₂), NHPh(2-NH₂), NHCH(CH₂CH(CH₃)₂)CO₂CH₂Ph, NHSO₂CH₂Ph(4-C(CH₃)₃), NHSO₂CH₂Ph, NHNHPh(2-F), NHCH₂Ph(4-CF₃), NHPh(4-OCH₂Ph), NHPh(4-SCH₃), NHCH(CH₂Ph)CO₂CH₂CH₃,
- NHCH(CH₂Ph)CO₂CH₃, NHCH₂Ph(4-OCH₃), NHCH₂-1-naphthyl, NHPh(4-F), NHCH₂Ph(2-F), NHCH₂CH(Ph)OH, NHCH₂CH₂Ph(4-F), NHC(CH₃)₂CH₂Ph(3-F), NHPh(3,4-diF), NHCH₂Ph(3-CH₃), NHNH(3-CH₃)Ph, NHCH₂Ph(2-Cl), NHCH₂Ph(2,4-diCl), NHNHPh(4-CH₃), NHCH₂Ph(4-Cl), NH(CH₂)₃Ph, NHCH₂CH₂Ph(4-Cl),
- 25 NHCH2CH2N(CH3)Ph, NHCH2Ph(3-CF3), NHCH2Ph(2-CF3), NH(CH2)4Ph, N(CH3)CH(CH3)CH(CH3)Ph, N(CH3)CH(CH3)Ph, N(CH2Ph)(CH2)3CH3, NHOCH2Ph, NCH2Ph(2,6-diF), N(CH3)CH(CH3)Ph, NHCH(CH3)Ph, N(CH3)CH2Ph, NHCH2Ph(3,4-diCl), N(CH3)CH(CH3)Ph,

N(CH₂Ph)CH₂CH₂Ph, NHNHCH₂Ph, NHCH₂Ph(2,4-diF), NHNHPh(2,5-diCl), NHCH₂Ph(3-F), NHCH(Ph)CH₂Ph, NHCH₂Ph(3,4-diOH), NHCH₂Ph(3,4-diOCH₃), N(CH₃)CH₂Ph, N(CH₂CH₃)CH₂Ph, N(CH₃)CH(CH₃)Ph, NHCH₂CH₂(3-F)Ph, NHCH(CH₂Ph)CH₂OH,

15. A compound having the formula

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wherein R8 and R9 are hydrogen and separated by a double bond or R8 is hydroxy, R9 is hydrogen and separated by a single bond and R^b is as defined in Claim 12.

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16. A compound having the formula

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wherein R8 and R9 are hydrogen and separated by a double bond or R8 is hydroxy, R9 is hydrogen and separated by a single bond and R^x is as defined in Claim 13.

17. A compound having the formula

- wherein R8 and R9 are hydrogen and separated by a double bond or R8 is hydroxy, R9 is hydrogen and separated by a single bond and NR^xRy is as defined in Claim 14.
 - 18. A compound having the formula

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wherein R^x is as defined in Claim 13.

19. A compound having the formula

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wherein NRXRY is as defined in Claim 14.

20. A compound having the formula

NHR*

wherein R^{χ} is as defined listed in Claim 13.

5

21. A compound having the formula

OH

OH

OH

OH

wherein NRXRY is as defined in Claim 14.

10

22. A compound having the formula

NH NHR'

OH OH

wherein R^X is as defined listed in Claim 13.

15

23. A compound having the formula

wherein NRXRY is as defined in Claim 14.

5 24. A compound having the formula

where R₁ - R₆, R₈ and R₉ are as defined in Claim 1;

R₁₁ is

(1) COCl,

10

- (2) CON₃, or
- (3) NCO.

25. A pharmaceutical composition comprising a compound of Claim 1 and a pharmaceutically acceptable carrier.

15

26. A composition of Claim 25 further comprising an anthelmintic agent.

20

27. A composition of Claim 26 wherein said anthelmintic agent is selected from ivermectin, avermectin, abamectin, emamectin, eprinamectin, doramectin,

fulladectin, moxidectin, Interceptor and nemadectin, thiabendazole, cambendazole, parbendazole, oxibendazole, mebendazole, flubendazole, fenbendazole, oxfendazole, albendazole, cyclobendazole, febantel, thiophanate, tetramisole-levamisole, butamisole, pyrantel, pamoate, aoxantel or morantel.

- 28. A composition of Claim 25 further comprising fipronil, lufenuron or an ecdosyne agonist.
- A method for treating a parasitic disease in a mammal which comprises administering to said mammal an antiparasitic effective amount of a compound of Claim 1.
- A method of Claim 29 further comprising administering an anthelmintic agent.
- 31. A method of Claim 29 further comprising administering fipronil or lufenuron.

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INTERNATIONAL SEARCH REPORT

In tional application No.
PCT/US96/03611

A. CLASSIFICATION OF SUBJECT MATTER IPC(6) :A61K 31/40, 31/425, 31/445, 31/495; C07D 405/06, 487/16 US CL :514/233.2, 255, 322, 365, 397; 544/142, 310; 546/199; 548/417 According to International Patent Classification (IPC) or to both national classification and IPC					
B. FIELDS SEARCHED					
Minimum documentation searched (classification system follow	wed by classification symbols)				
U.S. : 514/233.2, 255, 322, 365, 397; 544/142, 310; 546/199; 548/417					
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched					
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) CAS Structure					
C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category* Citation of document, with indication, where	appropriate, of the relevant passages Relevant to claim No.				
X, P US 5,399,582 A (A. W. DOMBI 1995, columns 1-2, compounds					
Further documents are listed in the continuation of Box	C. See patent family annex.				
Special categories of cited documents:	"T" later document published after the international filing date or priority				
'A' document defining the general state of the art which is not considered	date and not in conflict with the application but cited to understand the principle or theory underlying the invention				
to be of particular relevance E* earlier document published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be				
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O* document referring to an oral disclosure, use, exhibition or other means	combined with one or more other such documents, such combination being obvious to a person skilled in the art				
P [*] document published prior to the international filing date but later than the priority date claimed	*&* document member of the same patent family				
Date of the actual completion of the international search	Date of mailing of the international search report				
10 MAY 1996	20 MAY 1996				
Name and mailing address of the ISA/US	Authorized officer				
Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231	Jacqueline Haley Allon to				
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